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NUCLEIC ACIDS

STRUCTURES, PROPERTIES, AND FUNCTIONS

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Conformational Changes

CHAPTER

by Douglas H. Turner

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9. PREDICTING THREE DIMENSIONAL

STRUCTURE

OLIGONUCLEOTIDES WITH LOOPS

from Tetrahymena thermophila (Cech and Bass, 1986). At high temperatures, this molecule has the conformation of a random coil, and is not active. At temperatures formations. For example, Figure 8-1(a) shows the sequence of a self-splicing intron near 37°C, however, the interactions discussed in Chapter 7 force the molecule to fold on itself to give the base pairing, or secondary structure, shown in Figure 8-1(b) (Cech To be functional, nucleic acids must adopt particular three dimensional (3D) con-

аместиминастилизация силинастилиства весеми в серописти в серопист повети повети в серопист повети в серопист повети в серопист повети

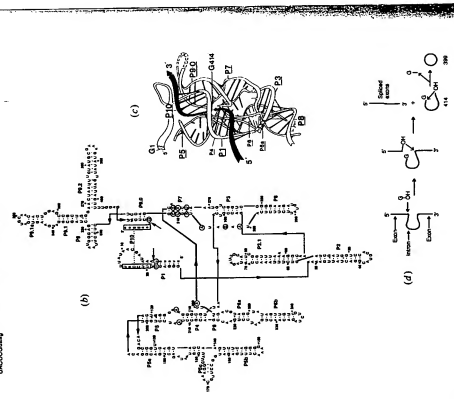


Figure 8-1 (facing page).

(a) The sequence of the self-splicing intron from the rRNA precursor of *T. thermophila* is shown in capital letters. Small letters give the adjacent sequences of the attached exons.

(b) Secondary structure formed by the sequence in (a). [Reprinted with permission from Cech, R. R., Damberger, S. H., and Gutell, R. R. (1994). *Nature Struct. Biol.* 1, 273–280.] Nucleotides that are circled are conserved in other group I introns. (c) Proposed model for 3D structure of active site for self-splining intron [Reprinted with permission from Michel and Westhof, 1990].

(d) Sequence of reactions mediated by structure in (b). [From Cech and Bass, 1986.]

et al., 1994). These same interactions also determine the 3D folding of the molecule, which has been suggested to look like Figure 8-1(c) at the active site (Michel and Westhof, 1990). When the molecule is in the necessary conformation, it can catalyze the reactions shown in Figure 8-1(d) (Cech and Bass, 1986; Cech, 1990). Since transcription of RNA occurs by stepwise addition of nucleotides, conformational changes are required for it to assume its active state.

Self-splicing introns are examples of nucleic acids with catalytic functions. The Self-splicing introns are examples of nucleic acid, however, is to carry information, such as the sequence of a protein. In these cases, conformation is still important for interactions with other molecules that regulate expression and transmission of the information (Oxender et al., 1998; Tang and Draper, 1989).

As described in Chapter 3, it is now straightforward to determine the sequence of a nucleic acid. Sequences containing more than 1 billion nucleotides are known, and determination of the 3 billion nucleotides of the human genome is in sight. Determination of the secondary and 3D structures and of the dynamics of these structures is more difficult, however (see Chapters 4 and 5). This finding is particularly true for RNA. Thus, transfer RNA is the only complete natural RNA whose 3D structure is known. While DNA overwhelmingly occurs as a fully paired, right-handed double helix, X-ray and solution studies reveal considerable sequence-dependent differences in conformation (Saenger, 1984; Dickerson, 1983). Presumably, these differences are important for interactions of DNA with other molecules. Thus the sequence dependence of conformation and its functional effects are important, but relatively unexplored.

One of the goals of biophysical chemistry is to use the available information on the structure and dynamics of nucleic acids to deduce principles that allow accurate prediction of these properties from sequence alone. In this chapter, experimental results no conformational changes are described, and the disciplines of thermodynamics, stalts on conformational changes are described, and the disciplines of thermodynamics, bused to test predictions made from our knowledge of intermolecular and intramolecular forces as discussed in Chapter 7. Knowledge of these forces will ultimately permit accurate predictions of properties from sequence.

I. SINGLE-STRAND STACKING

Much biological activity of nucleic acids depends on regions of single-stranded, unpaired nucleotides. This configuration maximizes the number of active groups available

Chapter 8 / Conformational Changes

of the codon depends on pairing with the single-stranded anticodon. Transfer RNA for recognition or reaction. For example, in tRNA (see Figs. 4-11-4-14), recognition is charged with an amino acid at one of the hydroxyl groups of the 3'-terminal A in the single-stranded sequence NCCA. As shown in Figure 8-1, several single stranded nucleotides of the self-splicing intron are conserved. Presumably they are important for catalysis (Kim and Cech, 1987; Michel and Westhof, 1990).

One of the simplest conformational changes in nucleic acids involves the transition of an unpaired single strand from a random coil in which the bases are not stacked to an ordered, helical structure in which the bases are stacked. This transition is illustrated in single strands. Most of the methods used in these studies are also applicable to other Figure 8-2. The thermodynamics and dynamics of this transition have been studied for a number of cases, and provide insight into the interactions governing the properties of conformational changes.

Fransition from unstacked to stacked conformation in single-stranded dinucleoside monophosphate.

1.1 Thermodynamics of Single Strand Stacking

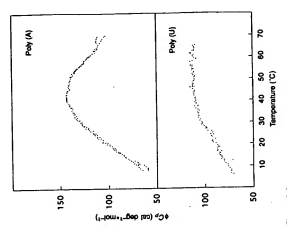
properties. Both methods are easy to understand in principle. In practice, neither is formational changes are calorimetry and the temperature dependence of spectroscopic easy to use with transitions for single-strand stacking because the transitions occur over a wide temperature range. For example, reported enthalpy changes, ∆H°, for The main methods for determining thermodynamic parameters for nucleic acid consingle strand stacking in poly (A) range from -3 to -13 kcal mol-1.

1.1.1 Calorimetry

of two matched cells. One cell contains the sample of interest and the other contains a blank (e.g., buffer). If the temperature change induces a chemical reaction in the sample but not the blank, then the amount of electrical energy required to raise the The most popular calorimetric method applied to conformational changes of nucleic 1992). In this method, electrical energy is used to slowly increase the temperatures temperature of the sample cell will be increased by the amount of heat absorbed by the reaction. Thus a differential scanning calorimeter measures the difference in heat Calorimetric methods measure the heat released or absorbed by a chemical reaction. acids is differential scanning calorimetry (DSC) (Sturtevant, 1987; Breslauer et al.,

required to raise the temperatures of the two cells. The data is reported as excess heat capacity, φC,, versus temperature as shown in Figure 8-3.

One disadvantage of DSC is that any reaction occurring in the sample cell can affect the excess heat capacity. For example, if hydration of the polymer chain is temperature dependent, then ϕC_{ρ} will be affected over the temperature range where hydration is changing. This disadvantage is not serious for studies of conformational changes that occur over small temperature ranges. In these cases, baselines before and after the transition of interest can be extrapolated to subtract out from ϕC_s any effects from other reactions. This subtraction leaves the change in heat capacity, ΔC_p^o , associated with the transition of interest. The area under the AC, curve for the temperature



Associated with the Conformational Transitions of Polyriboadenylic Acid and Polyribouridylic Acid. Suurkuusk, J., Alvarez, J., Ferier, E., and Biltonan, R., Biopolymers, 16, 2641–2652. Copyright © 1997. Reprinted by permission of John Wiley & Sons, Inc.] Excess heat capacity, ϕC_p , versus temperature for poly(A) and Calorimetric Determination of the Heat capacity Changes poly(U) as measured by differential scanning calorimetry. Figure 8-2

interval, T1 to T2, in which the reaction is occurring is the ΔH° for the transition (sec Figs. 8-3 and 8-4):

$$\Delta H^{\circ} = \int_{\gamma}^{T_{\rho}} \Delta C_{\rho}^{\circ} dT \tag{8-1}$$

in the normally accessible temperature range of 0-100°C. This limitation makes it Unfortunately, the stacking reaction for a single stranded polynucleotide is incomplete difficult to determine the baseline relevant for measuring area under the excess heat capacity curve.

1.1.2 Temperature Dependence of Spectroscopic Properties

The ΔH° for a reaction is related to the temperature dependence of the equilibrium constant by the van't Hoff equation:

$$\frac{\partial \ln K}{\partial T} = \frac{\Delta H^{\circ}}{RT^{2}} \tag{8-2}$$

Thus any method that provides an equilibrium constant as a function of temperature can be used to determine ΔH° . Spectroscopic methods are most common. For example, the equilibrium constant for the reaction shown in Figure 8-2 is

$$U \rightleftharpoons S \quad K = \frac{|S|}{|U|} = \frac{\alpha}{1 - \alpha} \tag{8.3}$$

property that allows determination of [S] and [U] can be used to determine K. For example, the extinction coefficients for stacked and unstacked conformations are usually different (see Chapter 6, Section 1.1). Thus the absorbance, A, of a sample containing where [S] and [U] are concentrations of stacked and unstacked species, respectively, and $\alpha = \{SI/(\{S\} + \{U\}), \text{ the fraction of strands in the stacked state. Therefore any$ dinucleoside monophosphate will be

$$A = \varepsilon_{s}[S]\ell + \varepsilon_{u}[U]\ell = C_{T}\ell[\alpha\varepsilon_{s} + (1 - \alpha)\varepsilon_{u}]$$
 (8-4)

concentration, $C_T = \{S\} + \{U\}$. Unfortunately, it is often difficult to determine ε , and e, particularly when the transition occurs over a large temperature range so that the respectively, and ℓ is the pathlength of the cell. If ϵ , and ϵ are known, then the relevant concentrations can be determined from the absorbance and knowledge of the total Here s, and s, are extinction coefficients for the stacked and unstacked species, sample is never completely one species.

8-2, the two-state model is the simplest and most common model. The dinucleoside When e, and e, cannot be directly measured, plots of absorbance versus temperature must be fit to a model for the transition of interest. For the transition shown in Figure monophosphate is assumed to be either stacked or unstacked, and the ΔH^a and ΔS^a

1 Single-Strand Stacking

for the reaction are assumed to be independent of temperature. For this model, plots of absorbance versus temperature can be fit with four variables: $\varepsilon_i, \varepsilon_s, \Delta H^*$, and ΔS^* ,

$$K = \exp(-\Delta H^{\circ}/RT + \Delta S^{\circ}/R)$$
 (8-5)

This treatment assumes the change in absorbance is entirely due to the stacking reaction and this assumption can be checked by measuring the time dependence of the absorbance change as described in Section 8.1.2.

Applequist, 1963). In this model, the equilibrium constants for propagating or initiating a stacked region are denoted s and β s, respectively. Thus β for a noncooperative transition is one. As a transition becomes more cooperative, β becomes smaller. While When single-strand stacking occurs in an oligonucleotide or polynucleotide, there coil. The equilibrium constants for these two processes can be different, indicating s has the temperature dependence given above for K, $oldsymbol{eta}$ is assumed to be temperature is an additional consideration. A new stack can either lengthen an existing region of stacked nucleotides, or it can occur in a region that was previously completely random cooperativity for stacking. In this case, a somewhat more complex model, the onedimensional (1D) Ising model, must be used for the analysis (Zimm and Bragg, 1959; independent. For this model, a is given by (Applequist, 1963):

$$\alpha = 0.5 + 0.5(s - 1)[(1 - s)^2 + 4\beta s]^{-1/2}$$
 (8-6)

The addition of $oldsymbol{eta}$ means five parameters are required to fit data to the model. In practice, the transitions are too broad to reliably allow a five parameter fit. Simultaneous fitting of spectroscopic and calorimetric data can improve the situation, however (Freier et al.,

Representative thermodynamic parameters for some dinucleoside monophosphates and polynucleotides are listed in Table 8-1. Included in the table are melting temperatures, t_n , the temperatures where half the bases are stacked (K=1 and/or $\alpha=0.5$).

$$\Delta G = -RT \ln K = \Delta H' - T\Delta S \tag{8-7a}$$

Celsius (°C). Thus, for a unimolecular transition, $U \rightleftharpoons S$, at the T_m , $-RT_m \ln(1) = 0 =$ Here T is the Kelvin temperature, T=273.15+t, where t is the temperature in degrees $\Delta H^{\circ} - T_{m} \Delta S^{\circ}$, so

$$T_{\rm m} = \frac{\Delta H^{\circ}}{\Delta S}$$
 (unimolecular transition) (8-7b)

 \deg^{-1}). Note that round off errors often affect T_m values by several degrees. While measured T_m 's are reasonably reliable, caution is required when considering values of ΔH° and ΔS° for single-strand stacking. As noted above, all involve extrapolation All temperatures in the above equations are in K. When the units of ΔH° are kilocalories per mole, they must be multiplied by 1000 if ΔS° is in entropy units (eu, cal mol-1

Э -1.0-10.0 01-0.6-UqU Simpkins and Richards (1967). A,ORD 1.0 No stacking 0 > CPC Powell et al. (1972). 1-10.0 **č.8−** 95-Olsthoom et al. (1981). UqA CD €.7-77 57-AqA Powell et al. (1972). ٧ 1-10.0 **c.8−** 87-97 AqA Olsthoom et al. (1981). CD 2.r-77 52-AqAb Olsthoom et al. (1981). CD €.T--53 6Þ Keterence MOIECUIE (N9CI) Method g (kcsi mol-1) (Gn), э. ۳, .5⊽ ۰н۷

Methods: A = absorbance: C = calonimetry: CD = circular dichroism: ORD = optical rotatory dispersion: S = sedimentation: V = viscosity.

S,V

D,A

D,A

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2.0

1.0-20.0

20.0

1

٥.5°

6.0

Richards et al. (1963); Inners and Felsenfeld (1970).

Thermodynamic Parameters for Single-Strand Stacking

Freier et al. (1981).

Freier et al. (1981).

Suurkuusk et al. (1977).

Filimonov and Privalov (1978).

of data outside experimentally accessible temperature limits. As discussed in Section 8.1.2, kinetic results indicate the two-state model also may not be adequate for certain

poly(C) are largely stacked at 20°C, poly (U) is a random coil. Structural studies of oligomers by NMR and ORD have provided evidence that the negligible stacking of U is not restricted to UU sequences. For example, in the sequences AUG and UGUG, the purines stack together while the U bases remain unstacked (Lee and Tinoco, 1980; Despite the uncertainties, some features of single-strand stacking are clear. The cooperativity is small, with $oldsymbol{eta}$ typically between 0.5 and 1. The salt dependence is also negligible. The sequence dependence, however, is considerable. While poly(A) and van der Hoogen et al., 1988b).

1.2 Kinetics of Single Strand Stacking

No stacking

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1.2.1 Temperature-Jump Relaxation Spectroscopy

Eq. 8-2). If the temperature-jump occurs faster than the equilibrium can adjust, then the time dependence of the approach to the new equilibrium concentrations can be ime dependence of the change in concentration of unstacked species, $\Delta[U]$, is a single which changes the equilibrium constant for any reaction that has a nonzero ΔH^{lpha} (see followed spectroscopically. The rate constants for the reaction can be derived from this time dependence. For example, for the two-state reaction shown in Figure 8-2, the 1976; Turner, 1986). In this method, the temperature of a solution is raised quickly, The kinetics of single strand stacking are very fast. They can be measured, however by temperature-jump relaxation spectroscopy (Eigen and De Maeyer, 1963; Bernasconi exponential:

$$\Delta[U] = \Delta[U]_0 \exp(-t/\tau) \tag{8-8}$$

jump, and i is time. For a unimolecular reaction, $\tau^{-1} = k_1 + k_{-1}$, the sum of the forward and reverse rate constants. The equilibrium constant is given by $K = k_1/k_{-1}$. Thus Here $\Delta\{U\}_n$ is the displacement from the new equilibrium at the time of the temperature. measurements of the relaxation time, r, and K allow determination of k, and k_1.

1.2.2 Results

poly(A) (Freier et al., 1981; Pörschke, 1978; Dewey and Turner, 1979). These results there is more than one conformational change in these cases. Studies by NMR indicate this results because two stacked conformations exist (Kondo and Danyluk, 1976). Thus Single relaxation times have been measured for CpC, CpA, ApC, poly(C), and are listed in Table 8-2. At Na * concentrations above 0.05 M, the relaxation times for poly(A) and ApA depend on the monitoring wavelength (Pörschke, 1978), which means

Average of two values.

Poly(U)

Poly(C)

Poly(A)

Poly(A)

Poly(A)

All the forward rate constants for single strand stacking are about 10° s-1. While this value is large compared to many other reactions, it is slow for a simple stacking reaction. For example, the relaxation time for stacking of the two adenines joined by the two-state model is not appropriate, and rate constants cannot be derived easily.

Molecule	Reference	(M ₈ +1)	1 D°	(su)	اد- ^s لا ^ا × 10و	(kcsl mol ₋₁)	(na) ¹S∇	°-1 1'-1 × 10e	(kcal mol ⁻¹)	(na) ¹-¯S∇
AqA	Pörschke (1978).		<i>b</i>	50, > 100	<i>D</i>			U		
⊃q∧	Pörschke (1978).	I	Þ	45	13.			. II		
AqD	Pörschke (1978).	1	Þ	30	. † I			,61		•
OqO	Pörschke (1978).	I	Þ	30	,61	1.2	61-	, † I	Ħ	71
^9—(CH ₂),—•A	Pörschke (1978).	I	Þ	\$1 >			-	,.	٧.	01
– (A)yloq	Dewey and Tumer (1979).	\$0.0	01	\$LI	2.9	7.9	S.T-	9 [.] I	₹i	81
Poly(C)	Freier et al. (1981).	\$0.0	01	St	61	8.0-	0ε-	1.5	8.T	z-
Poly(C)	Freier et al. (1981)	Ţ	۶I	<i>L</i> 9	٤١	6.1	-33	8.1	11	9
(Ab)ylo4	Dewey and Turner (1979).	20.0	10	81	77	€.€	ςı-	4.5	9.6	ε

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1 Single-Strand Stacking

268

are provided by the activation energies, E_o , and entropies, ΔS^4 , as derived from the in a single-stranded polymer, the magnitude of the unimolecular rate constant would be at least as large if controlled by the same factors. These comparisons suggest the sugar-phosphate backbone somehow limits the stacking rate. Further insights into this 1978), indicating $k_{\rm i} > 7 \times 10^3 {\rm s}^{-1}$. The forward rate constant for bimolecular stacking reactions of bases and of planar dye molecules is typically about 109M-1s-1 (Pörschke and Eggers, 1972; Dewey et al., 1979). Given the high local concentration of bases a trimethylene bridge in 9,9'-trimethylenebisadenine is faster than 15 ns (Pörschke, temperature dependence of rate constants and the Eyring equation:

$$k = (eRT/\hbar N) \exp(-E_g/RT + \Delta S^t/R)$$
 (8)

sion controlled rate. The activation entropies, however, are about -10 to -30 cu. Thus ble contributor to the large, unfavorable ΔS^4 is specific solvation requirements for the Here e is the base for natural logarithms (2.72), h is Planck's constant, and N is Avogadro's number. For the stacking of poly(C) and poly(A), E, is only a few kilocalories per mole (Freier et al., 1981; Dewey and Tumer, 1979). Furthermore, stacking rate constants depend linearly on solvent viscosity. This finding is consistent with a diffuthe relatively slow rate constants are a consequence of a very ordered transition state, which could result from the necessity of constraining bonds in the backbone to single conformations before rotational diffusion to the stacked conformation. Another possitransition state.

1.3 Interactions Determining Structure and Dynamics of Single-Strand Stacking

tions, in order to stabilize the ordered, stacked helix. Both effects are discussed below, sight into the interactions that determine these properties. In particular, conformational entropy favors the unstacked, random coil conformation. These unfavorable entropy The results for the thermodynamics and kinetics of single-strand stacking provide ineffects must be overcome by bonding interactions, commonly called stacking interacalong with some unexplained effects.

1.3.1 Conformational Entropy Effects

available conformations for the bonds shown in Figure 2-3 are (Berman, 1981; Olson, 1982): $\alpha(1)$, $\beta(3)$, $\gamma(3)$, $\delta(2)$, $\varepsilon(3)$, $\zeta(2)$, $\chi(2)$. If the stacked state of a dinucleoside monophosphate is restricted to a single conformation, but the unstacked state can sample all available conformations for each bond, then initiation of stacking will be opposed by an unfavorable initiation conformational entropy term of For most bonds in polymers, the number of available conformations is typically three. X-ray and NMR data on nucleic acids suggest the appropriate number of There are a limited number of conformations that can be adopted around each bond. The covalent bonds in a dinucleoside monophosphate are shown in Figure 2-3.

entropy change, ΔS_{κ} , of -11 eu. The values measured for ΔS^{κ} and ΔS^{t} of stacking in small cooperativity observed. If the cooperativity parameter $oldsymbol{eta}$ arises only because two glycosidic and two sugar bonds are constrained for initiation and one each for propagation, then its value is predicted to be $\beta=\exp[(-R \ln 2 \times 2)/R]=0.25$. Measured values range from about 0.5 to 1. Of course, this model may be oversimplified. For dinucleoside monophosphates and in polymers range from about - 10 to -30 eu (see Tables 8-1 and 8-2). Thus conformational effects can account for a large part of the measured equilibrium and activation entropy changes. They can also account for the cosidic (χ) and two sugar (δ) bonds are included in this calculation, one for each ing an adjacent nucleotide on the pre-existing stack would require constraining rotation about one more glycosidic and one more sugar bond, rather than two of each. In this simple model, propagation of a helix is associated with a propagation conformational = $-R \ln(1 \times 3 \times 3 \times 2 \times 2 \times 3 \times 2 \times 2 \times 2) = -13$ eu. Notice that two glynucleoside. If the stacked dimer in Figure 8-2 was part of a polynucleotide, then stackexample, more than one configuration may contribute to the stacked state.

1.3.2 Stacking

The origin of stacking interactions has been the subject of much discussion. From the thermodynamic data, it is clear that stacking is driven by a favorable, modest ΔH° . This finding is consistent with noncovalent bonding due to dispersion forces as described in Section 7.1.

strand stacking, however, is unfavorable (see Table 8-1). Moreover, the magnitude is It has often been suggested that stacking is also driven by classical hydrophobic interactions. In this model, ordered water is released from around the bases upon stacking, and this provides a favorable entropy term. The magnitude expected for such an effect can be estimated from the dimerization of benzene in water. The $\Delta\mathcal{S}$ for benzene dimerization is a favorable 20 eu (Tucker et al., 1981). The AS for single either that expected for conformational entropy effects, or is even more unfavorable. Thus the thermodynamic parameters provide no evidence that classical hydrophobic bonding is important for driving stacking.

1.3.3 Unexplained Effects

in poly(C) and poly(A) are also suggestive of a difference (Table 8.2). The $E_{a,1}$ and ΔS_1^i for poly(A) are reasonable for a diffusion controlled process. For poly(C), however, $U \leq C < A$ versus UpU \ll CpC < ApA versus poly(U) \ll poly(A) < poly(C). This result suggests some new interaction in poly(C). The activation parameters for stacking $E_{a,1}$ is lower and ΔS_1^4 higher than expected for diffusion control. One possibility is a The results on single strand stacking also raise some interesting questions. Comparison of Tables 8-1 and 2-2 indicates the ΔH° values for stacking of ApA, poly(A), and deoxyadenosine monomers are similar. The ΔH° values for stacking of CpC and poly(C), however, are very different from the ΔH^a for association of cytidines. Moreover, the order for strength of stacking in monomers, dimers, and polymers is different: special, specific solvation of poly(C).

2 Double Helix Formation by Oligonucleotides without Loops

2. DOUBLE HELIX FORMATION BY OLIGONUCLEOTIDES WITHOUT LOOPS

Crick base pairs. Cellular DNA is almost exclusively in this form. Known structures of RNA are more than 50% double helix. Thus an understanding of the principles governing double helix formation is essential for understanding and predicting the properties of nucleic acids. Oligonucleotides provide convenient model systems for The main structural motif for natural nucleic acids is the double helix with Watsondiscovering these principles.

2.1 Thermodynamics of Duplex Formation

2.1.1 Methods

The methods used for measuring the thermodynamics of double helix formation formation, however, is a very cooperative process, so the transitions occur in smaller temperature intervals. For example, Figure 8-4 shows a DSC curve for duplex formation by oligonucleotides are similar to those used for single-strand stacking. Double helix

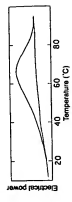


Figure 8-3

Reprinted with permission from Albergo, D. D., Marky, L. A., Breslauer, K. J., and Turner, D. cacodylate, pH 7. For the lower curve, both sample and reference cells contained only buffer. sample cell to maintain it at same temperature as reference cell) versus temperature. For the upper curve, the sample cell contained $1\times 10^{-3}\,M\,{\rm dGCGGG}$ in 1 M NaCl, 45 mM Raw data from a DSC experiment on dGCGCGC. Shown is electrical power (fed back to H. (1981). Biochemistry, 20, 1409-1413. Copyright @American Chemical Society.]

baselines outside the transition region are apparent, which makes integration of the ΔC_{ρ} curve to obtain ΔH° relatively reliable (see Eq. 8-1). To obtain ΔS , the same by dGCGCGC (Albergo et al., 1981). The beginning and end of the transition and the data can be plotted as $\Delta C_{\sigma}/T$ versus temperature and integrated:

$$\Delta S = \int_{1}^{7} \frac{\Delta C_{p}}{T} dT \tag{8-10}$$

An advantage of calorimetry is that the thermodynamic parameters obtained in this way do not depend on a theoretical model for the transition. As long as the baselines can be determined, the thermodynamic parameters can be obtained by taking the area under the transition curve.

The temperature dependence of spectroscopic properties is also used to obtain thermodynamic parameters for duplex formation. In this case, the data must be fit to a theoretical model to derive parameters. In practice, a simple two-state model is used most often (Martin et al., 1971; Borer et al., 1974; Turner et al., 1988; Petersheim and Turner, 1983). The simplifying assumption in the two-state case is that a given strand is either maximally base paired or completely not base paired (see Fig. 8-5). This assumption corresponds to a completely cooperative transition. More general statistical mechanical treatments are discussed in Appendix 8-11.

The equations used to fit spectroscopic data to the two-state model depend on whether the oligonucleotides are self- or nonself-complementary:

Self-complementary
$$2A \rightleftharpoons A_2$$
 $K = \frac{[A_2]}{[A]^2} = \frac{\alpha}{2(1-\alpha)^2 C_T}$ (8-11)

Nonself-complementary
$$A+B \rightleftharpoons AB$$
 $K = \frac{[AB]}{[A|[B]} = \frac{2\alpha}{(1-\alpha)^2 C_T}$ (8-12)

Here α is the fraction of total strand concentration, C_T , that is in duplex; and for the nonself-complementary case, it has been assumed that the total concentration of A and B strands is each $C_T/2$. If the spectroscopic property is absorbance, A, then at any

Self-complementary
$$A = C_T \ell[\epsilon_A (1 - \alpha) + \epsilon_{A_1} \alpha/2]$$
 (8-13)

Nonself-complementary
$$A = \frac{C_1 \ell}{2} [(\epsilon_{\lambda} + \epsilon_{B})(1 - \alpha) + \epsilon_{AB} \alpha]$$
 (8-14)

Here ϵ_{λ} , ϵ_{B} , $\epsilon_{\Lambda_{\lambda}}$ and ϵ_{AB} are extinction coefficients for single-stranded A and B, and for duplexes A_{2} and AB, respectively. If these extinction coefficients are known, then absorbance versus temperature data can be fit with Eqs. 8-11-8-14 and 8-7a to provide values of ΔH^{α} and ΔS^{α} for the transition. In practice, the extinction coefficients are usually temperature dependent. For example, the extinction for a single strand will vary



Figure 8-4 Transition from two single strands to a double helix.

with temperature because of the stacked-to-unstacked equilibrium discussed in Section 8.1. When nonself-complementary oligomers are studied, this temperature dependence can sometimes be measured independently with the individual single strands. In most cases, the temperature dependences of extinction coefficients are assumed to be linear, for example, $s_A = m_A T + b_A$. The values for m_A and b_A are determined by fitting absorbance versus temperature data in the regions where only single strands or only duplexes occur, or by including the linear dependence of extinction in fitting the shape of the entire duplex-to-single strand transition curve. Note that not every spectroscopic property is as simple as absorbance. For example, NMR chemical shifts cannot easily be used to derive thermodynamic properties (Pardi et al., 1981).

For a two-state transition, the concentration dependence of duplex formation provides another method for determining thermodynamic parameters. Defining the melting ing temperature, T_m , as the temperature at which $\alpha=0.5$ and plugging into Eqs. 8-11 and 8-12 gives the results that at the T_m , K is L/C_m for a self-complementary transition and $4/C_m$ for a nonself-complementary transition. Substituting these results into $\Delta C = -RT \ln K = \Delta H^* - T \Delta S^*$, and retarranging leads to

Bimolecular, self-complementary
$$\frac{1}{T_m} = \frac{R \ln C_T}{\Delta H^*} + \frac{\Delta S^*}{\Delta H^*}$$
 (8-15a)

Bimolecular nonself-complementary i
$$R \ln(C_T/4) + \Delta S$$
 (8-15b) $\frac{1}{T_m} = \frac{R \ln(C_T/4)}{\Delta H^o} + \frac{\Delta S}{\Delta H^o}$

Thus, a plot of $1/T_m$ versus $\ln(C_{+})$ should be linear, and ΔH^n and ΔS^n can be determined from the slope and intercept. Equations 8-15a and 8-15b are special cases of the general equations for N strands associating to form an N-mer (Marky and Breslauer, 1987):

N-mer, Self-complementary
$$\frac{1}{T_m} = \frac{(N-1)R}{\Delta H^{\sigma}} \ln C_{\tau} + \frac{[\Delta S^{\sigma} - (N-1)R \ln 2 + R \ln N]}{\Delta H}$$
(8-16a)

N-mer, Nonself-complementary
$$\frac{1}{T_n} = \frac{(N-1)R}{\Delta H^n} \ln C_T + \frac{\lfloor \Delta S^n - (N-1)R \ln 2N \rfloor}{\Delta H^n}$$
(A.16a)

If a transition is two state, then all the methods described above should give the same thermodynamic parameters. If a transition is not two state, then the magnitude of ΔH^a determined from calorimetry will be larger than that determined spectroscopically (Sturtevant, 1987). Moreover, the two spectroscopic methods may give different results, since non-two-state behavior typically affects the shape of a melting curve more than the T_m . These comparisons, therefore, provide tests for two-state behavior.

2.1.2 Results for Duplexes without Loops

Table 8-3 lists thermodynamic parameters determined for duplex formation by some oligonucleotides. The enthalpy and entropy changes are quite large relative to

respectively (Martin and Tinoco, 1980). It has been suggested that the relative stabilities of RNA. RNA, DNA. DNA, and RNA. DNA duplexes may determine the positions har have been studied, duplexes with antiparallel strands are more stable than duplexes 1992; Jin et al., 1993). For 3'-5'linked duplexes, whether a deoxy or ribo duplex is more stable depends on the sequence. Different stabilities might be expected for sequences containing AU or AT pairs. The RNA and DNA sequences with only GC pairs also have different stabilities, however. Thus the 2'-OH affects duplex stability. This 2'-OH effect is even observed if a single 2'-OH is replaced by H in an oligonucleotide (Bevilacqua and Turner, 1991). This effect may be related to the different conformational preferences for RNA and DNA (see Chapter 4). For RNA-DNA hybrid duplexes, the available data suggest interesting sequence dependence (Martin and Tinoco, 1980; Sugimoto et al., 1995). For example, at total strand concentrations of $4 \times 10^{-4} M$, the melting temperatures of rCA3G-dCT3G and dCA2G-rCU3G are 19 and less than 0°C. oxy and ribo forms, in parallel and antiparallel strands, and in 3'-5'and 2'-5'linked oligomers. The thermodynamic parameters are different in each case. For sequences with parallel strands (Rippe and Jovin, 1992), and duplexes with 3'-5'phosphodiester bonds are more stable than duplexes with 2'-5'phosphodiester bonds (Kierzek et al., Table 8-3 contains some results for sequences that have been studied in both de-

for factor-independent termination of transcription (Martin and Tinoco, 1980).

Table 8-3 also illustrates the large dependence of stability on base pair composition.

At 10⁻⁴M oligonucleotide strands, the duplex with 9 bp formed by CA,G and CU,G melts at 32°C, whereas the 4 bp duplex (GCCC), melts at 34°C. Evidently, GC pairs are more stable than AU pairs. The sequence dependence of stability depends on more than composition, however. For example, another duplex with four GC pairs, (CGCG)₂, malts at hown 19°C at 10⁻⁴M.

neighbors and base compositions but different sequences indicates the thermodynamic Nearest neighbor models (Borer et al., 1974; Turner et al., 1988; Goldstein and tions and the base composition of the helix as reflected by the terminal base pairs and 5'GG3'/3'CC5' are different. Thus (GGCC), and (CGCG), are composed of different nearest neighbors and can have different stabilities in the model. The duplexes (GCCGGC), and (GGCGCC), however, both have the same nearest neighbors and same base compositions, and therefore must have identical stabilities within the model. Inspection of the experimental results in Table 8-3 for GCCGGC ($t_m = 67.2^{\circ}$ C) and GGCGCC (1, = 65.2°C) indicates the nearest neighbor model is a reasonable approximation for this case. A study of pairs of oligonucleotides with identical nearest parameters for such oligomers are generally within about 10% of each other (Kierzek et al., 1986; Xia et al., 1998). Thus the nearest neighbor model is better than a simple reasonable approximations of the sequence dependence of duplex stability. One such model, the Independent Nearest Neighbor-Hydrogen Bonding or INN-HB model, assumes that the stability of a given base pair depends on the identity of the adjacent base pair and that the stability of a helix depends on these nearest neighbor interac-Benight, 1992; Gray 1997a,b; Allawi and SantaLucia, 1997; Xia et al., 1998) provide (Xia et al., 1998). For example, the nearest neighbors 5'CG3'/3'GC5', 5'GC3'/3'CG5', composition model, but is not perfect. melts at about 19°C at 10-4M.

The 8.3 Oligonucleotides for Duplex Formation by Oligonucleotides

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(D°) (°	\(\rho \text{Cosl mol}\) -4.4 -3.4 -3.4 -3.4 -3.4 -11.2	(u9) \$\sigma \cdot \cdo	-28°4 -28°1 -28°1 -38°1 -3°8 -3°8 -3°8 -3°8 -3°8	(D°) 27.1 26.5 36.4 26.5 36.4 27.1 19 27.1	AC37 (kcal mol-1) -4.6 -3.7 -4.6 -5.4 -9.1 -7.4	7:111- 1:86- 1:86- 9:56- 9:56- 9:56-	(kcal mol ⁻¹) -34.2 -34.2 -34.2 -54.5 -54.5 -54.5	Reference 1 M Na.Cl Freis et al. (1998). Freis et al. (1998). Freis et al. (1983). Freis et al. (1983). Freis et al. (1983). Freis et al. (1983). Freis et al. (1984).	RNA, 3'-5', Antiparallel Strands, CCCG CCCG
8.81	6.E- 7.A- A.E- A.E- A.T-	2.59- 4.501- 5.201- 8.821- 2.721-	8.25- 8.35- 1.85- 1.82- 1.82-	91 8.65 9.46	1.6- \$.2- \$.4- 7.6- \$.4-	6.26- 6.26- 6.28- 1.89- 1.89-	2.AE- 6.6E- 2.0E- 8.2E- 8.2E-	I. M Na.C.] Petersbeim and Turner (1983). Xia et al. (1998). Freier et al. (1983). Freier et al. (1983). Freier et al. (1983). Freier et al. (1983).	RAA, 3'-5', Antiparallel Surands. CCGG CGCG GCGC GCGCG
8.81	6.E- 7.A- A.E- A.E- A.T-	2.59- 4.501- 5.201- 8.821- 2.721-	8.25- 8.35- 1.85- 1.82- 1.82-	91 8.65 9.46	7.E- 6.A- 4.2- 1.9-	6.26- 6.28- 1.89- 6.641-	6.65- 2.05- 8.25- 8.45-	Peterabeim and Turner (1983). Xis et al. (1998). Freier et al. (1985b). Freier et al. (1985b). Groebe, Cameron. Freier, Turner, and	939393 9399 9393 9393 9393
	p.2- p.2- b.4-	2.59- 4.501- 5.201- 8.821- 2.721-	8.25- 8.35- 1.85- 1.82- 1.82-	91 8.65 9.46	7.E- 6.A- 4.2- 1.9-	6.26- 6.28- 1.89- 6.641-	6.65- 2.05- 8.25- 8.45-	Xis et al. (1998). Freier et al. (1985b). Freier et al. (1986b). Groebe, Cameron. Freier. Turner, and	939393 3399 3939 9393
1.62	4.6- 6.7-	6.601- 2.201- 8.621- 2.721-	8.85- 1.85- 1.82- 1.82-	26.5 94.4 9.72	6.4- 4.2- 1.9-	p.68- 1.89- p.341-	2.05- 8.25- 2.42-	freier et al. (1985b). Freier et al. (1985). Freier et al. (1986b). Groebe, Cameron, Freier, Tumer, and	939393 2399 2939
	4.6- 6.7-	8.821 - 2.721 -	1.82- 1.82- 4.62-	6.72	4.2- 1.9-	1.89-	8.2E- 2.42-	Freier et al. (1983). Freier et al. (1986b). Groebe, Cameron, Freier, Tumer, and	939393 3399
94.9	9.T-	2.721-	1.82- 4.32-	6.72	1.6-	7'971-	2.42-	Freier et al. (1986b). Groebe, Cameron, Freier, Turner, and	939393
2.82								Groebe, Cameron, Freier, Tumer, and	
48.3	2.11-	8.891-	919-					Ubleabeck, unpublished results.	PCVDPC
9.99									
1.58	2.01 -	0.781-	6.£8- E.£8-	2.73	2.11-	0.661-	F.53-	Freier et al. (1985b).	299229
9.99	2.11-	8.891-	9.69-	1.29	9.01-	2.871-	0.99-	Freier et al. (1985b).	292929
4.SE	T.2-	2.691-	2.82-	2.23	ביוו-	0.281 -	8.75-	Freier et al. (1986b).	229299
			7:05-	9.15	5.2-	1.271-	8.66-	Nelsen et al. (1981); Freier et al. (1986b).	c∧,G + CU,G
					• •				RNA. 2'-5', Antiparallel Strands.
				24.4	8.1-	2.95-	7.22-	Kierzek et al. (1992).	292929
				45.3	8.9-	2.121-	E. DD-	Kierzek et al. (1992).	9000000
۲ >>	70	,	•.,						DNA, 3'-5', Antiparallel Strands
7.22 2.8£	0.8-	4.751-	2:15-	T.ZZ	€.8-	8.221-	4.94-	Senior et al., (1988b).	
2.82		9.121~	8.Eb-	£.8£	6.5-	811-	42.4	Williams et al. (1989.	333333
1.54		2. PEI -	\$.05~	1.62	1.6-	L'791-	9.65-	Albergo et al. (1981).	GCATGC
1:55	8.9-	5.281-	5.59-	1.05	Z.T.	961-	0.89-	Aboulels et al. (1985)	0.00000
									CA,G+CT,G DNA, 3'-5', Antiparallel Strands
						LSV-	151-	Rippe and Jovin (1992); Rentzeperis (1992).	`č.TAT.A,T.AT.O.A'?
						- 200	<i>L</i> 91-	Rippe and lovin (1992); Rentzeperis (1992).	'c, ATAD, T, ADTAD, AD, T'S
	2.9-	PSE-	911-			,,,,			DNA, 3'-5', Parallel Strands, 0.1 S' A ₁₀ , TA ₇ , T ₄ , TAT ₉ , 3'
			011			7 18-	911-	Rippe and Jovin (1992); Rentzeperis (1992).	'ε, λτλ, τ, λ, τλ" Τ'ς 'ε, τλτο, λ, τολτο, λο, Λ'ς
						186-	911-	Rippe and Iovin (1992); Rentzeperis (1992).	s'T,CATCA,T,CATA,3'

 $\pi = \Lambda H' - 310$ 14.5.7 may differ from the listed $\Delta G_{\rm c}$, due to round off errors. The $c_{\rm c}$ is for 1×10^{-3}

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hose for single-strand stacking. In contrast to single-strand stacking, these changes also increase substantially in magnitude as the sequence is made longer (e.g. GCCGGC

Sugimoto et al., 1995). For example, at total strand concentrations of $4 \times 10^{-4} M$, the respectively (Martin and Tinoco, 1980). It has been suggested that the relative stabilities of RNA · RNA · DNA · DNA, and RNA · DNA duplexes may determine the positions 1992; Jin et al., 1993). For 3'-S'linked duplexes, whether a deoxy or ribo duplex is more stable depends on the sequence. Different stabilities might be expected for sequences containing AU or AT pairs. The RNA and DNA sequences with only GC pairs also have different stabilities, however. Thus the 2'-OH affects duplex stability. This 2'-OH effect is even observed if a single 2'-OH is replaced by H in an oligonucleotide (Bevilacqua and Turner, 1991). This effect may be related to the different conformational preferences for RNA and DNA (see Chapter 4). For RNA-DNA hybrid duplexes. the available data suggest interesting sequence dependence (Martin and Tinoco, 1980; melting temperatures of rCA, G·dCT, G and dCA, G·rCU, G are 19 and less than 0°C, oxy and ribo forms, in parallel and antiparallel strands, and in 3'-5'and 2'-5'linked oligomers. The thermodynamic parameters are different in each case. For sequences that have been studied, duplexes with antiparallel strands are more stable than duplexes with parallel strands (Rippe and Jovin, 1992), and duplexes with 3'-5'phosphodiester Table 8-3 contains some results for sequences that have been studied in both debonds are more stable than duplexes with 2'-5'phosphodiester bonds (Kierzek et al., ys. CCGG). These trends reflect the high cooperativity of the duplex transition. for factor-independent termination of transcription (Martin and Tinoco, 1980).

are more stable than AU pairs. The sequence dependence of stability depends on more At 10-4M oligonucleotide strands, the duplex with 9 bp formed by CA,G and CU,G melts at 32°C, whereas the 4 bp duplex (GGCC), melts at 34°C. Evidently, GC pairs than composition, however. For example, another duplex with four GC pairs, (CGCG),, Table 8-3 also illustrates the large dependence of stability on base pair composition. melts at about 19° C at $10^{-4}M$.

neighbors and base compositions but different sequences indicates the thermodynamic parameters for such oligomers are generally within about 10% of each other (Kierzek et al., 1986; Xia et al., 1998). Thus the nearest neighbor model is better than a simple tions and the base composition of the helix as reflected by the terminal base pairs and 5'GG3'/3'CC5' are different. Thus (GGCC), and (CGCG), are composed of diferent nearest neighbors and can have different stabilities in the model. The duplexes (GCCGGC), and (GGCGCC), however, both have the same nearest neighbors and nspection of the experimental results in Table 8-3 for GCCGGC ($t_m = 67.2^{\circ}$ C) and = 65.2°C) indicates the nearest neighbor model is a reasonable approximation for this case. A study of pairs of oligonucleotides with identical nearest Nearest neighbor models (Borer et al., 1974; Turner et al., 1988; Goldstein and reasonable approximations of the sequence dependence of duplex stability. One such sumes that the stability of a given base pair depends on the identity of the adjacent same base compositions, and therefore must have identical stabilities within the model. Benight, 1992; Gray 1997a,b; Allawi and SantaLucia, 1997; Xia et al., 1998) provide model, the Independent Nearest Neighbor-Hydrogen Bonding or INN-HB model, asbase pair and that the stability of a helix depends on these nearest neighbor interac-(Xia et al., 1998). For example, the nearest neighbors 5'CG3'/3'GC5', 5'GC3'/3'CG5' composition model, but is not perfect. ") ככככ (v"

Thermodynamic Parameters for Duplex Formation by Oligonucleotides

Petersheim and Turn Xia et al. (1998). Freier et al. (1985b)

Keletence

and Turner (1985).

RNA, 3'-5', Antiparallel Strands, 1 M NaCl

						185-	911-	Rippe and Jovin (1992); Rentzeperis (1992).	יג'סאלסלגס, איסינאלי, מזאז, ז' ז'ד, כאזאס, ד'באזאל, ד'נ
	2.9-	-324	911-			PLE-	911-	Rippe and lovin (1992); Remixeperis (1992).	`£,TAT,A,T,AT ₀₁ A`2 `£,ATA,T,A ₅ TA ₀₁ T`2
						006-	L91-	Rippe and Iovin (1992); Rentzeperis (1992).	5'A,GA,GTAGT,A,GTAT,3' 3'T,CG,CATCA,T,CATA,5' DNA, 3'-5', Parallel Strands, 0.
						LST-	151-	Rippe and Jovin (1992); Rentzeperis (1992).	`£ _e TAT _e A¸T _e AT _{es} A`è `è _e ATA _e T _e A _e TA _e ,T'€
								1.0.1 M NaCi	DNA, 3'-5', Antiparallel Strand
						961-	0.89-	Aboutela et al. (1985)	ο,τ⊃+ο,∧⊃
Þ	8.9-	2.281-	2.69-	1.05	2.7-	7.231 -	9.65-	Albergo et al. (1981).	202020
\$	T.8-	6.4€! —	4.02-	1.88	6.2- 1.9-	811-	4.54-	Williams et al. (1989.	OOTA DO
Ē	6.8-	9.121-	ð.£4—	€.8€	£.8-	8.221-	4.34-	Senior et al., (1988b).	900000
;	6.8 -	1.751-	2.12-	r.ee	£ 8-	8	• ,.	I N N*CI	NA, 3'-5', Antiparallel Strands
				#1#s	0:0 -	2.121-	£.44	Kierzek et al. (1992).	93393993
				45.3	8.4— 8.4—	2.95-	2777	Kierzek et al. (1992).	292929
				24.4	8 V-	6 93		DIN WIL	2.14, 2-5', Antiparallel Strands
					e-e -	1.271-	8.66-	Nelsen et al. (1981); Freier et al. (1986b).	5A.G + CU,G
ε	T.2-	2.691 -	2.88-	9.15	¿.ç.	0.281-	8.73-	Freier et al. (1986b).	229299
•	2.11-	8.891-	6.£9 −	2.23	£.11-	2.871-	0.55-	Freier et al. (1985b).	202029
•	201-	0.731 -	£.53-	1.29	2.11- 2.01-	0.881-	L'79-	Freier et al. (1985b).	202229
•	2.11-	8.831-	6.53 -	£.73	211-	0.791	,	Upleabeck, unpublished results.	:cvnec
	0:1 -	2.721-	4.05-	L'SV	p.Y-	T:LL1-	6.23-	Groebe, Cameron, Freier, Turner, and	939393
۶ ۲	4.6- 6.7-	8.321-	1.82-	6.72	1'6-	p'971-	242-	Freier et al. (1986b).	30308
		0 731	1.85-	34.4	P.C-	1.86-	8.25-	Freier et al. (1983).	2020

2.45-2.55-2.05-8.25-2.42-5.42-

(kcsl mol_,)

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6.29-6.29-1.89-1.89-1.81-

(Ga)

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8.81 1.92 1.94.9 8.82 8.82

(aⁿ)

0.29-2.29-4.501-2.201-8.321-2.721-

(na)

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8.25-8.25-8.85-1.85-1.82-

(kcsl mol_)

1.72 2.62 34.4 6.72 7.24

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6.6-6.6-6.6-1.6-6.7-

(kcal mol

Measured

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6.6-6.6-6.8-6.8-6.7-

(KCS) WOL

Predicted

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8.4 Phole Brameters for Helix Initiation and Propagation in IM NaCI

-132 -13 -132 -13 -134 -13 -137 -13 -137 -13 -137 -13 -137 -13 -137 -13 -137 -13	-8.0 -12.8 -12.8	\(\text{C3}\) \(\text{(kcsl mol-1)}\) \(\text{-2.17}\) \(\text{-2.17}\)	2.12- 2.12- 6.61- 4.42- (na) •2.5∆	ΔH° (kcsl mol ⁻¹) -9.8 -8.0 -10.6	∆C ₂ , (kcal mol ⁻¹) -3.26 -3.26 -2.36	.25.7 -36.9 -32.7 -32.7	(kcal mol ⁻¹) -14.88 -14.88 -14.88	PAV	CG C
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61- 616-	8:21- 6:91-	₽8.1 ~ C1.5 ~	6.61-	0.8 8.01	65.5-	7.2£-	66.81 - 48.01 -	GA	70 50 50 50 50 50 50 50 50 50 50 50 50 50
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¿.1- e.ss-								10	വാ
	9.8-							-	-
1.1- 2.15-	8.7-	** .1-	4.22-	4.8-	-2.24	£.95.	04.11-	± 10	on →
1.2- 6.51-	6.6-							→ CV	→ C¥
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	,VNQ.VN	В		DNY,			KNY,			
(kcal mol⁻¹)	(na) .5∇	(kcsl mol_1) ∇H。	(kcal mol−¹) ∆C₃,	(en) \$₹	(kcsl mol_1) ∇H。	(kcal mol-³)	(en) ∇2.	(kcsl mol_₁) ∇H。		noisgation eaneupe
6.0-	£-61-	0.7-	82.1-	0.15~	8.7-	80.5-	1.72-	89:01-	<u> </u>	na -
8.1-	2.55-	1.6-							 VD	→ v o
9.0-	2.55.	8.7-	82.0-	£.12-	2.1-	££.1-	\$.02-	69°L-	← ΛΤ	v ∩
									ıv →	nv →
6.0-	6.55-	£.8-	88.0-	4.05-	2.r-	01.1-	T.82-	86.9-	← TA AT	UA AU
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			\$0.0	6'9	77.	54.0	2.1- 2.01	Σ <i>Γ.</i> Ε	TA to UA	Initiation Each terminal
			64.0	* I=	0	64.0	p.1 -	0	вопов	умтиену сок (вец-сошрієни
			0	0	0	0	0	0		тоэ ультау сот (ровзей-сотр

Yis et al., (1998).
*Allawi and SantaLucia (1997).
*Sugmono et al., (1995). For RMA/DMA hybrids with two sets of parameters, the top set corresponds to the top strand as RMA and the bottom set corresponds to the bottom set RMA/DMA data base have also been surmed as RMA. For example, APP \$'GCG' = -12.8 letal mol⁻¹ and APP \$'GCG' = -9.3 letal mol⁻¹. Alternative analyses of the RMA/DMA data base have also been presented (Gray, 1997b).

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analysis has also been done for the available data on parallel stranded double helixes Application of the INN-HB nearest neighbor model to RNA and DNA duplexes Gray, 1997a,b). The parameters have been determined by fitting to optical melting data for oligonucleotides (Xia et al., 1998; Allawi and SantaLucia, 1997; Sugimoto predictions from the model with measurements are shown in Table 8-3. For DNA, (Breslauer et al., 1986). SantaLucia (1998) and Owczarzy et al. (1997) compare the containing only Watson-Crick pairs requires determining thermodynamic parameters for 10 nearest neighbors, helix initiation, and helix termination by AU or AT for each case. For RNA/DNA hybrids, more parameters are required (Sugimoto et al., 1995; et al., 1995; Gray, 1997b). Some results are listed in Table 8-4, and comparisons of calorimetric data for both oligomers and polymers have also been fit to a similar model thermodynamic parameters for DNA obtained by various methods. A nearest neighbor (Rippe and Jovin, 1992).

or DNA duplex containing only Watson-Crick base pairs. The melting temperature in degrees Celsius, $I_{\mathbf{m}}$, for any concentration can then be calculated from a rearrangement In deriving the parameters in Table 8-4, it was necessary to consider the fact that whereas single-strands and nonself-complementary oligomers do not. To correct for this symmetry difference, AS for self-complementary oligomers must be reduced by R1n2 = 1.4 cu. Including this correction and simply summing up appropriate parameters allows prediction of thermodynamic parameters for the melting of any RNA duplexes formed by self-complementary oligomers have a twofold rotational symmetry, of Eqs. 8-15a and 8-15b:

Bimolecular, self-complementary
$$t_{\rm m} = \frac{\Delta H^{\circ}}{\Delta S^{\circ} + R \ln(C_{\rm T})} - 273.15$$
 (8-17)

Bimolecular nonself-complementary
$$I_m = \frac{\Delta H^o}{\Delta S^o + R \ln(C_T/4)} - 273.15$$
 (8-18a)

Equation 8-18a holds if the concentrations of the two nonself-complementary strands are equal. In many applications, including probing with oligonucleotides for complementary sequences in polynucleotides, one strand is in large excess. In this case, when $_{\rm m}$ is defined as the temperature where half of the less concentrated sequence is bound:

Bimolecular nonself-complementary
$$t_m = \frac{\Delta H^*}{\Delta S^* + R \ln(C_B - 0.5C_A^*)} - 273.15$$
(8-18b)

Equations 8-17 and 8-18a can be combined into one by including all the changes between self- and nonself-complementary oligomers in a constant, A:

$$I_{\rm m} = \frac{\Delta H^{\circ}}{A + \Delta S_{\rm NN} + R \ln(C_{\tau})} - 273.15$$
 (8-19)

oligomers, respectively. Note that in Eqs. 8-17-8-19, if the units for ΔH° are in Here, ΔS_{NN} is the entropy change without any symmetry term, C_{τ} is always the total strand concentration, and A is -1.4 and -2.8 cu for self- and nonself-complementary

kilocalories per mole, they must be multiplied by 1000 if AS is in entropy units.

Sample calculations are shown in Figure 8-6.

interactions of the 3'A with the opposite strand help hold the duplex together, whereas for DNA, but it appears that 5'-dangling ends will add more stability than 3'-dangling can stabilize a helix more than some base pairs (cf. Tables 8-5 and 8-4), indicating a opposite strand G of the adjacent base pair, the S'-dangling A of (AGGCC), is not close to the opposite strand C of the adjacent base pair. Thus it is not surprising that interactions of the S'A with the opposite strand are negligible. Fewer data are available helix (Freier et al., 1985). Stability increments for 3' dangling ends are sequence dependent, however, ranging from -0.1 to -1.7 kcal mol-1. Thus some 3' dangling ends strong interaction. This difference between 5' and 3' stacking can be rationalized from in A- form geometry. Whereas the 3'-dangling A of (GGCCA), stacks directly on the The available parameters are listed in Table 8-5 (Turner et al., 1988). One striking averaging -0.2 kcal mol-1. Similar free energy increments are measured for addition of a S'-phosphate suggesting that S'-dangling ends interact little with the adjacent structural considerations. Figure 8-7 shows stercoviews of (AGGCC)₂ and (GGCCA)₂ The results in Table 8-4 allow prediction of the thermodynamic properties of fully paired duplexes. Many nucleic acid associations, however, include additional sequence dependent (Turner et al., 1988). For example, at 10-4 M strands, the melting temperatures of (GGCC), (AGGCC), and (GGCCA), are 34.4, 38.1, and 57.9°C, respectively (Freier et al., 1983b, 1985a). The effects of at least the first unpaired nucleotide adjacent to a helix can be approximated by a nearest neighbor model. observation for RNA oligomers in 1 M NaCl is that all the free energy increments for terminal unpaired nucleotides on the 5' side of helixes are very similar and small, unpaired nucleotides on the ends of double helixes. Two examples are the associations of IRNA and mRNA and of hybrid probes with DNA or RNA. For RNA, these unpaired nucleotides or "dangling ends" add stability to the double helix in a manner that is very ends (Senior et al., 1988b; Mellema et al., 1984).

for interactions between the bases in most terminal mismatches. The exceptions are A 5' and 3' dangling end opposite each other is called a terminal mismatch. Free energy increments associated with terminal mismatches are listed in Table 8-6 (Freier et al., 1986a; Hickey and Turner, 1985a; Sugimoto et al., 1987b; Serra et al., 1994). For pyrimidine-pyrimidine and CA mismatches, these stability increments are essentially the sum of the increments for the constituent dangling ends. For purine-purine mismatches, the increment may be less. Thus there is no thermodynamic evidence terminal GU mismatches. In certain cases, the pairing of a terminal G and U provides more stability than the sum of the equivalent dangling ends. This finding is consistent with hydrogen-bond formation within GU mismatches as suggested by Crick (1966) in the "wobble" hypothesis (see Chapter 2).

2.2 Interactions Determining Stability of the Double Helix

Conformational entropy and stacking interactions present in single-stranded nucleic acids are also present in duplexes. In addition, hydrogen bonds are formed between base pairs, and duplex formation leads to increased condensation of counterions around

6.9

7.0

E.E

2.2

L'0-

8.5-

1.4-

6.0-

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877

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6°L

8.1-

6.T-

L'01-

2.1-

.5∇

າ=x

£.0-

7.0-

2.0-

€.0+

L'0-

1.1-

L'1-

8.0-

"ე⊽

Table 8.5 Thermodynamic Parameters for Unpaired Terminal Mucleotides in RNA at 1 M NaCl'

2.08 -2.36 -2 5'	in + 2AGTERM-AU
(a) 5' AGCGCU + + + + + + + + + + + + + + + + + + +	$\Delta G_{\text{true}}^2 + \Delta G_{\text{true}}^2 + \Delta G_{\text{true}}^2 + \sum \Delta G_{\text{true}}^2 + 2\Delta G_{\text{true}}^2 - \Delta G_{\text{true}}^2$

$$\Delta G_{\rm BOT}^2 = \Delta G_{\rm BOT}^2 + \Delta G_{\rm SYM}^2 + \Sigma \Delta G_{\rm SNI}^2 + 2\Delta G_{\rm TEMM}^2$$

= 409+0.43+(-1336)+2×0.45

(c)
$$S'GUGUAAUAACC3' \stackrel{3.26}{\longleftarrow} 1 1 1 A \stackrel{A}{\longleftarrow} 1 1 A \stackrel{A}{\longleftarrow} 1 A$$

 $\Delta G_{TOT}^{\circ} = \Delta G_{INIT}^{\circ}(n=6) + \Delta G^{\circ}\left(\mathrm{First\ Mismatch}\right) + \Delta G^{\circ}\left(\mathrm{Sum}\right)$ = 5.4 + (-1.1 - 0.8) + (-5.05) = -1.55 kcal/mol

Figure 8-5 Preficing stability with the nearest neighbor model of Xia et al. (1998). (a) Calculation of free energy change for duptex formation by a self-complementary sequence. (b) Calculation for energy change for duptex formation by a self-complementary sequence. (a) Calculation for nonself-complementary sequence. Calculations of ΔH^* and ΔS^* are similar to those for ΔG^* , except there is no symmetry term for ΔH^* . Thus for the top sequence, $\Delta H^* = 3.61 + 2(-10.84) + 2(-11.88) + (-10.64) + 2(3.72) = -50.31 \, kcal mol⁻¹, <math>\Delta S^* = -1.5 - 1.4 + 2(-27.1) + 2(-36.9) + (-25.7) + 2(10.5) = -13.66 \, cu, and t_a at 10^{-4} M strands = 324.8 K = 51.67. Cr the sequence in (b), <math>\Delta H^* = 3.61 + (-11.40) + (-10.64) + (-12.44) + (-10.48) + (-14.88) + 3.72 = -23.51 \, kcal mol⁻¹, <math>\Delta S^* = -1.5 + (-29.5) + (-25.5.7) + (-25.7) + (-25.7) + (-25.7) + (-25.7) + (-25.7) + (-25.7) + (-27.7) + (-26.7) + ($

3, Unpaired Nucleotides

Sedneuce

Propagation

2.0-

9.1-

77-

9.I

r.e-

p. L-

0.6-

∘н⊽

1.8

p:91-

0.02-

4.22-

7.61-

۵۶⊽

∀=X

2.08 ↓ 3′ ↓ U ↑ 5′ 1.045

=-7.94 kcal/mol

(よな)

8.41-

3.4

Þ.E

p:91-

9.91-

2.22-

0.21-

۰5⊽

y=c

(z.0-)

0.0

2.0-

p.0-

T.O-

£1-

L1-

8.0-

"ე⊽

(L.S)

2.1-

£.4-

9.01

L'6-

⊅.0Z-

4.2-

.5⊽

n=x

(2.0-)

0.0

1.0-

Z:0-

1.0~

9.0~

z:1-

9.0-

∜D⊽

(9.0)

Þ.O.

1.5

7.2-

9.£−

£.2-

∘н⊽

(0.0)

8.0

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1.0-

8.0-

¿.0-

450₹

= 4.09 + 0 + (-12.45) + 0.45 =-7.91 kcal/mol

Table 8.6 Thermodynamic Parameters for Terminal Mismatches in RNA*

	U -0.8	U -0.5 -0.5
	C C C C C C C C C C C C C C C C C C C	
ΔG³, (kcal mol ⁻¹)	U X/Y A (-0.5) C (-1.0) (-0.7) U -1.4	0.77 A -1.0 (-0.7) C -0.7 (-0.8) U -1.1 (-0.8) U -1.1
٥	10 10 10 10 10 10 10 10 10 10 10 10 10 1	G (-0.8)
	< 7.7.7.1 < 7.7.7.1	XX XX C C C C C C C C C C C C C C C C C

	U -2.7 -8.6	J 1.8 4.1+	
	-5.56 -9.26	28. 9.89. 9.99.	15
	28. €. 1999† €. 1993	2.66.22. 4.4.54.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4	
(,_	A 9.1 -8.2	4 4 1 1	
ΔH° (kcal mol '')		(-0.3) C (-1.7) U	
ρΉ	U .6 (-4.2) .2 (-5.0)		
	C C C C C C C C C C C C C C C C C C C	C G (+2.0) (-3.5) (+4.6) (-3.5)	t
	-5.2 -5.2 -7.2 -7.1 -7.1 -7.1 -7.1	-3.9 (+)	
	×<005	Ž<∪∪⊃	1
		0.0	-

	U -6.3 -23.9	U -4.2	+6.0
1561	C G -13.5 -13.4 (-7.6) -24.6 -12.6	UX AY AY -17.7 -23.2 -14.6 -25.0	
-	A -24.5 (-15.2)	- A -9.7 -11.6 -8.5	
	Ž _{∢∪©⊃}	۶۲۰۵	>
	U (-12.2) (-14.0)	U (+1.5)	(-2.7)
† 8 51	G 13.9 15.1	AAX UV + G (-8.7)	
1001	C (-8.2) (+3.9) (+2.1)		(+17.4)
	A -13.2 -19.6 -17.8	-10.2 -5.3 -7.3	
	Š<005	```	5

Freier et al. (1986a); Hickey and Turner, (1985); Sugimoto et al. (1987b); Serra et al. (1994). Values in parentheses are estimated.

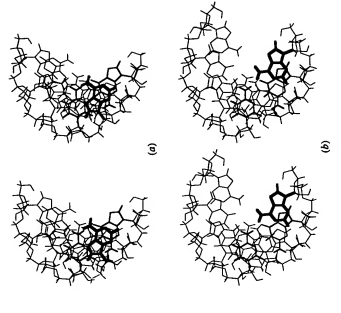


Figure 8-6 Stereoviews of (GGCCA), (a) and (AGGCC), (b) in A-form geometry. The terminal A closest to the reader is in boldface.

the backbone. All these factors affect the stability of the duplex. Solvent effects may also play a role. While exact partitioning of these effects on stability is not yet possible, a qualitative picture is emerging, and is discussed below.

2.2.1 Conformational Entropy

In Section 1.3.1, the conformational entropy associated with propagating a single-strand stacked helix by one additional nucleotide was estimated as -11 eu. Since propagation of a double helix by an additional base pair requires limiting the conformations accessible to two nucleotides, the conformational entropy associated with this process is estimated as $2 \times -11 = -22$ eu. Inspection of Table 8-4 indicates that 3/4 of the measured ΔS values for duplex propagation are within 6 eu of this theoretical

value. As with single strands, this suggests conformational entropy effects account for a large part of the observed ΔS

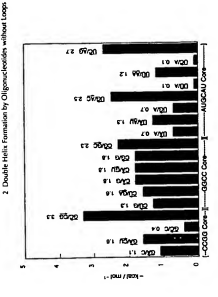
daralingam, 1969, 1973). This restriction could provide the small stabilization that is duplexes by 5'-dangling ends. As noted above, stabilization from a 5'-dangling end is about the same as from a 5'-phosphate. It has been suggested by Sundaralingam that addition of a 5'-phosphate restricts the conformations of the ribose group (Sun-Conformational entropy effects may also account for the stabilization of RNA observed

2.2.2 Stacking

may indicate no stacking at all for these sequences. This sequence dependence is A consistent with the idea that electronic interactions between the bases are responsible provided by the stability increments for 3'-dangling ends in Table 8-5. Comparison of some selected increments with those for base pairs is provided in Figure 8-8. This most 3'-dangling ends includes the unfavorable conformational entropy associated with that stacking. This unfavorable component of the stacking ΔG has been empirically considerations given above. When the empirical estimate of conformational effects is factored out, the favorable stacking interactions are estimated to be as large as -1.7 - 1.9 = -3.6 kcal mol⁻¹. The magnitudes of these favorable interactions are An empirical indication of the contributions of stacking to duplex stability is comparison is deceptive when considering the favorable attractive forces of stacking interactions, however, because the favorable free energy increment for stacking of estimated at about 1.9 kcal mol-1 at 37°C (Freier et al., 1986c), somewhat less than the value of $-7\Delta S = -(-11 \times 310) = 3.4$ kcal mol⁻¹ expected from the theoretical very sequence dependent. In fact, the negligible ΔG° values measured for UU or stacking.

2.2.3 Hydrogen Bonding

mol⁻¹. Stacking of each 3'-dangling end in (CCGGC), is associated with a ΔG_{31} of conformational entropy effects. For example (Fig. 8-8), in (GCCGGC), the $\Delta G_{\rm 37}^{\circ}$ for adding each terminal GC pair is $[\Delta G_{yy}^{\sigma}(GCCGGC) - \Delta G_{yy}^{\sigma}(CCGG)]/2 = -3.3$ kcal Inspection of Figure 8-8 indicates the stability increments for some base pairs are much larger than the increments for stacking of the constituent nucleotides. This suggests pairing between the bases contributes an additional interaction (Freier et hydrogen bonding can be made by taking stability increments for base pairs, subtracting stability increments for stacking of the constituent nucleotides, and correcting for al., 1986c; 1985a). Consistent with this are results with DNA oligomers showing that substitution of weakly hydrogen bonding diflourotoluene for T in an AT pair decreases duplex stability by 3.6 kcal mol-1 at 25°C (Moran et al., 1997). Presumably, the effect of hydrogen bonds is due to the difference between hydrogen bonding in a base pair and hydrogen bonding of the separated bases with water. Specific solvation effects could also be important. Estimates for the contributions of differential



Free energy increments associated with duplex formation at 37°C from $\binom{\Gamma^{0,0}}{2} = 1/2 \lfloor \Delta G_{j,j}^*(CCGGA) - \Delta G_{j,j}^*(CCGG) \rfloor = -1.1 \text{ kcal mol}^{-1}$ Note that the free energy increments for adding S' unpaired terminal GGCC, or AUGCAU cores. For example, the first bar on the left is ΔG_3 , for adding a 3' unpaired A adjacent to a GC pair, ΔG_3 , adding a terminal base pair or 3' unpaired terminal nucleotide to nucleotides are not shown but average only -0.2 kcal mol-1 Figure 8-7

G is $[\Delta G_{11}^{*}(GCCGG) - \Delta G_{12}^{*}(CCGG)]/2 = -0.2 \text{ kcal mol}^{-1}$ (This is no more than the G is not stacked.) Constraining the 5'-terminal G in a base pair, however, requires overcoming the estimated 1.9 kcal mol-1 of conformational free energy. Thus the free energy gained from pairing the terminal G and C is estimated as $\Delta G_{\rm Ho} = -3.3$ $(-0.4 - 0.2 + 1.9) = -4.6 \text{ kcal mol}^{-1}$. Since this pairing involves three hydrogen hydrogen bond. Similar calculations on other sequences give ΔG° values for a hydrogen Turner et al., 1987). It has been suggested that this range is due to a sequence dependent $\Delta G_{yy}(CCGGC) - \Delta G_{yy}(CCGG)]/2 = -0.4 \text{ kcal mol}^{-1}$. The effect of the 5'-terminal contribution from adding a 5' terminal phosphate suggesting an unpaired 5'-terminal bonds, the estimated free energy increment per hydrogen bond is -1.5 kcal mol-1 bond that range from -0.5 to -1.5 kcal mol-1 hydrogen bond (Freier et al., 1986c; competition between hydrogen bonding and stacking (Turner et al., 1987).

comparing duplex stabilities for sequences with different numbers of hydrogen bonds (Freier et al., 1986c; Turner et al., 1987). Such comparisons must also consider changes in stacking interactions that may result from changing the number of hydrogen bonding ing of 3'-terminal unpaired bases, there is little effect of changing hydrogen-bonding Another empirical estimate for the contributions of hydrogen bonds is provided by groups on the bases. Inspection of Table 8-5, however, indicates that at least for stack-

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with single changes in structure. For example, removing the 2-amino group from G comparisons of oligomers with similar sequences, but different numbers of hydrogen GC pairs are probably a second-order effect resulting from redistribution of electrons on pairing. Even these effects can be minimized by comparing stabilities of base pairs in a GC pair leaves an IC pair with two hydrogen bonds. Table 8-7 contains several U and C are also similar. Thus any large changes in stacking energies between AU and groups. Both A and G have very similar thermodynamic parameters for 3' stacking, and bonds. The stability increments per hydrogen bond range from -0.5 to $-2~\mathrm{kcal}$ mol $^{-1}$

2.2.4 Counterion Condensation

Repulsions of negatively charged backbone phosphates destabilize double helixes. Condensation of positively charged counterions around the helix favors duplex stability by neutralizing much of the negative charge on the phosphates. Experimental and theoretical aspects of this effect are discussed in Section 8.5, and in Chapter 11, respectively.

7

7

7

7

Z

Z

7

7

7

No. of H Bonds

Differences in

(5861

Turner et al. (1987)

Turner et al. (1987)

7.81

18.4

5.41

1.02

6.8

8.2

2.8

6.21

10~M(°C)

16 "1 <u>A</u> A

6.1-

8.1-

2.0-

2.0-

L'0-

9.1-

(kcal mol-1 H Bond)

Dad H\%D∆∆

2.2.5 Solvent

stacking. As discussed above, in most cases, the AS of duplex formation is similar to interactions are also associated with a large change in heat capacity, ΔC_{s} . Values of ΔC_p reported for single- to double-strand transitions of nucleic acids, however, are either modest or zero (Suurkuusk et al., 1977; Freier et al., 1983b, 1985a, 1986a,c; Kierzek et al., 1986; Breslauer et al., 1986). These generalizations even hold for duplexes containing thymine. Since thymine has a methyl group, some hydrophobic By analogy to proteins, classical hydrophobic interactions are often invoked as the view that classical hydrophobic interactions stabilize duplexes or single-strand or more unfavorable than expected from conformational terms. There is no indication of the large, positive ΔS expected for classical hydrophobic interactions. Hydrophobic a source of stability for double helixes. There is no evidence, however, to support interaction might be expected.

This topic is discussed in detail in Chapters 4, 7, and 11. Such specific solvation One solvent effect that may be important for helix stability is specific solvation. could explain the excess unfavorable ΔS observed for some helix transitions. The quantitative consequences of such solvation have not been determined, however.

2.3 Kinetics of Duplex Formation

2.3.1 Experimental Measurements

with the temperature-jump method. For the reaction shown in Figure 8-5, the time The kinetics of duplex formation by oligonucleotides have been measured primarily dependence of the change in concentration of single strands has the same form as Eq. 8-8. For duplex formation, the relaxation time, r, is given by

(8-21)(8-20) $\tau^{-1} = k_{\infty}([A] + [B]) + k_{\text{off}}$ $\tau^{-1} = 4k_{\rm on}[A] + k_{\rm off}$ Self-complementary Nonself-complementary

"I is inosine. A' is 2-aminoadenine.

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Freier et al. (1986b)	AUGUACAU	AUGCGCAU	
Freier et al. (1986b)	AUACGUAU	AUGCGCAU	
Freier et al. (1986b)	AAUGCAUU	GAUGCAUC	
Freier et al. (1986b)	NGCGCA	500000	
Freier et al. (1986b)	SCAUGC	292929	
Freier et al. (1986b)	GUGCAC	ეეეეეე	
Freier et al. (1986b)	cyecne	໑ͻͻ໑໑ͻ	
Gaffney et al. (1984)	ACGTACG	ACGTA'CG	
Kawase et al. (1986)	dG,A,ICT,C,	dG,A,GCT,C,	
Aboul-ela et al. (1982	CA,CA,G +dCT,IT,G	aca,ca,G+act,GT,G	
Martin et al. (1985)	acA,1A,G+acT,CT,G	aca,Ga,G+act,CT,G	
Turner et al. (1987)	CCCCI	໑ͻͻϧϧͻ	

ICCCCCC

Fewer H Bonds

Comparison of Stabilities of Duplexes with Different Numbers of Hydrogen Bonds'

sequence is self-or nonself-complementary. The activation energies for k_m and $k_{\rm eff}$ can be determined from measurements as a function of temperature (see Eq. 8-9). Here [A] and [B] are the equilibrium concentrations of the single strands at the higher temperature, $k_{\rm ca}$ and $k_{\rm caf}$ are the forward and reverse rate constants at the higher temperature as illustrated in Figure 8-5. Thus a plot of r-1 versus single-strand concentration has an intercept of k_{aff} and a slope of 4k_{an} or k_{an} depending on whether the

zero. That is, k can decrease with increasing temperature. Any elementary reaction step must have a positive activation energy. For example, diffusion controlled reactions in solution have activation energies of about 4 keal mol-1. Thus the mechanism of Typical results for the kinetics of duplex formation are compiled in Table 8-8. The forward rates, k_{α} , range from about $10^5-10^7\,M^{-1}s^{-1}$, and depend on both sequence and association of two like-charged species. The forward rate, however, is slower than the diffusion limit. Moreover, in several cases, the activation energy for k_m is negative or salt concentration. Higher salt concentrations increase the forward rate as expected for duplex formation must be more complicated than indicated in Figure 8-5.

is expected to be similar to that for single strand stacking, roughly 5 kcal mol-1. An the measured activation energy is approximated. For example, the measured activation measured activation energy for $k_{\rm m}$ depends on the $\Delta H^{\rm a}$ associated with formation of the intermediate preceding the nucleus and the activation energy for forming the last base pair in the nucleus (see Fig. 8-10): $E_{A,\infty} = \Delta H_{n-1}^{-1} + E_{A,n-1-n}$. Here n is the number of base pairs in the nucleus. The enthalpy change, ΔH_{n-1}^{-1} can be estimated from the values in Table 8-4. The activation energy for propagating the helix an additional base pair estimate of the size of the nucleus can be made by adding appropriate parameters until energy for duplex formation by A₆U₆ is -3 kcal mol-1 (Craig et al., 1971). Formation (Pörschke and Eigen, 1971; Craig et al., 1971; Williams et al., 1989; Wetmur and Davidson, 1968; Manning, 1976). In this mechanism, the rate-determining step is formation of a nucleus containing a small number of base pairs. This nucleus adds an additional base pair faster than it dissociates. Thus the double helix can "zip up" after formation of the nucleus. The stability of the intermediate preceding the nucleus, however, is temperature dependent. Like any double helix, its stability decreases as temperature increases. Thus the concentration of the intermediate preceding the nucleus decreases as temperature increases, and this can lead to a decrease in kan. The Figure 8-9 illustrates a mechanism consistent with the kinetic experiments

Mechanism for duplex formation. Step 0 is alignment of strands without inter-strand bonding. Subsequent steps involve formation of base pairs by hydrogen bonding. Figure 8-8

dGCATGC(0.01 M Mg ²⁺)	Williams et al. (1989).	210.0	1.15	EL	77	£-	EÞ.
		ı	1.15	6.6	2.2	ş-	07
		240.0	1.15	9.1	1.2	•	OV.
OSTAGE	Williams et al. (1989).	210.0	1.16	86.0	6.6	\$1	
OCCCAGANTICGCG	Chu and Tinoco (1983).	£1.0	8.15	70.0	7.0	91	LV
OCCIGAATTCGCG	Chu and Tinoco (1983).	41.0	8.15	80.0	0.1	7Z	7 L
•Tb + •A	Hoggen and Masss (1971).		23	01	740		89
J93939P	Freier et al. (1983a).	i	57	0.21	16.0	7-	£43
2229 + 2999	Podder (1971).	1.0	5.15	4.2		8.0	LS
4CA,G + 4CT,G	Nelson and Tinoco (1982).	,,	07		07	9.4	62
cv'e + cn'e	Nelson and Tinoco (1982).	;	1.12	6	04	2.0-	Er
$v^2c^3+c^3n^2$	Porschite et al. (1973).	80.0	£.ES	9.4	330	0	6ξ
v'œ¹ + c¹n¹	Pörschke et al. (1973).	20.0		4.4	340	L	EP
	Pörzchke et al. (1973).	500	8.91	7.11	350	٤١	9€
v'ecn'	Porschic et al. (1973).	1	23	6.0	ı	6	9 Z
v'ecn'	Porschic et al. (1973).	20.0	23.3	61.0	5.1	8	97
1100	Pörschke et al. (1973).	20.0	23.3	\$7.0	٤	L	05
v ³ ecn ³	Pörschke et al. (1973).	I	23	10	00Z	8	33
125		20.0	£.E.	9.1	057	٤	33
,υ,^A	Breslauer and Bina-Stein (1977).	ι	1.22	L'T	3.0	9-	50
"∩° v	Craig et al. (1971).	22.0	17	8.0	1	ς+	\$9
,;; ∧ , ∪, , ∧	Craig et al. (1971).	25.0	12	51	\$1	€-	09
'.''\ V'n'	Craig et al. (1971).	22.0	17	7	000Z	b -	05
*'n + *'\	Craig et al. (1971).	22.0	12	ı	3000	9-	LE
"U+"A	Pörschke and Eigen (1971).	20.0	23.5	19.0		8-	
	Pörschke and Eigen (1971).	20.0	23.3	05.0	017	zi-	ES
•0 + •v	Pörschlte and Eigen (1971).	\$0.0	€.£2	€8.0	7300	8-	9£
edneuce	Reference	[↑5N] (M)	J.	(10°M"s-1)	(s ₋₁)	(kcal mol ⁻¹)	(kcal mol ⁻¹)

Kinetic Parameters for Duplex Formation by Oligonucleotides 8.8 əldaT

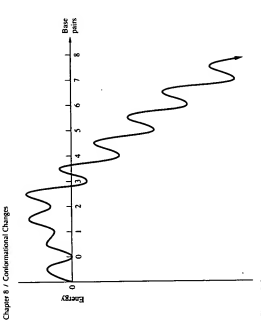


Figure 8-9 Energy profile corresponding to mechanism in Figure 8-9 for formation of duplex from separated single strands.

of four AU base pairs is associated with a $\Delta H^{\circ} \approx 3.6 + 2(3.7) + 3(-6.8) = -9.4$ kcal mol⁻¹. If addition of the next base pair requires an activation energy of 5 kcal mol⁻¹, then the overall activation energy would be $E_{\lambda,\infty} = -9.4 + 5 = -4.4$ kcal mol⁻¹. Thus the results are consistent with a nucleus of 5 bp. Since several assumptions are required, the estimate for the size of the nucleus is rough. In general, the results in Table 8-8 are consistent with nuclei of 5 ± 1 bp for obligomers containing only AU pairs, and 2 ± 1 bp for obligomers containing at least two GC pairs.

The data in Table 8-8 can also be used to estimate a rate constant of about 10⁷ s⁻¹ for addition of the next base pair to the nucleus (Pörschke and Eigen, 1971; Craig et al., 1971). This rate constant is similar to that for single-strand stacking, which supports the choice of activation energy used above.

The activation energies reported for homo-duplex formation by dCGTGAATTCGCG and dCGCAGAATTCGCG are larger than expected from the above analysis (Chu and Tinoco, 1983). Both these oligomers are long, self-complementary, and contain non-Watson-Crick pairings. One possibility is that each oligomer has intramolecular base pairs that must be broken before duplexes can form. Another is that non-Watson-Crick pairings may introduce effects that are not easily predicted from available information. Further experiments are required to sort out these effects.

3 Double Helix Formation by Oligonucleotides with Loops

These results are consistent with the model described above. Dissociation of duplex requires breaking enough base pairs to return to the nucleus. The energy required for this will depend on sequence and length. If the size of the nucleus and the activation energy for losing a base pair from the nucleus are known, then the ΔH^a values in Table 8.4 can be used to predict $E_{A,art}$. For example, for A_bU_a , 7 bp must be broken to return to the 5 bp uncleus. The activation energy for the reverse rate is predicted to be $E_{A,art} = 6 \times 6.8 + 9.4 + (5 + 6.8) = 62$ kcal mol⁻¹. The measured value is 60 kcal mol⁻¹ (Craig et al., 1971).

2.3.2 Predicting Kinetics of Duplex Formation

The results described above suggest a way to roughly predict the kinetics of duplex formation between oligonucleotides that do not have intramolecular base pairing. At high Na² concentration or in the presence of Mg²⁺, the forward rate is almost always $10^{M}H^{-1}s^{-1}$ within an order of magnitude, and the temperature dependence is modest. The equilibrium constant for duplex formation can be predicted from the parameters in Table 8-4, since $K = \exp(-\Delta G'/RT)$ and $\Delta G' = \Delta H' - T \Delta S$. The predicted values of K and k_{cc} can then be used to predict k_{cc} since $K = k_{cc}H'_{cc}$. The predicted values of K and 3.1×10^{12} , respectively, giving predicted off rate constants of 1.5 × 10² and 3×10^{-3} s⁻¹, respectively, The half lives for dissociation of these helixes are predicted from $t_{t/t} = \ln(2)/k_{cd} = 0.693/k_{cd}$ to be about S ms and 27 days, respectively, at $37^{\circ}C$. Once that the off rate is very temperature dependent. For example, at $0^{\circ}C$, the half lives for (GGCC) and (GGCCGCCC), are predicted to be about 20 s and 30 million years. This type of vough approximation is often sufficient for designing experiments.

3. DOUBLE HELIX FORMATION BY OLIGONUCLEOTIDES WITH LOOPS

Loops are regions of non Watson-Crick paired nucleotides flanked by one or more double helixes. The known varieties of loops are illustrated in Figure 8-11. All occur in structures of RNA (e.g., see Fig. 8-1). While natural DNA is primarily double helix, loops can occur transiently or even be favored under certain conditions. Thus it is important to understand the properties of loops.

There are several experimental ways to determine the thermodynamic properties of loops. All involve measuring properties for structures containing the loop of interest and subtracting out other contributions. For example, ΔG for the bulge loop in the duplex formed by GCG Δ GCG + CGCCGC is obtained by taking ΔG for this bulge duplex and subtracting ΔG for the fully paired duplex formed by GCGGCG + CGCCGC:

 $\Delta G_{\text{bulk}} = \Delta G^{*}(\text{GCG}\underline{\text{A}}\text{GCG} + \text{CGCCGC}) - \Delta G^{*}(\text{GCGGCG} + \text{CGCCGC})$

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The assumption is that the loop does not affect the stabilities of other regions so the free energies are additive. This assumption may not be very good (Longfellow et al., 1990). Often loops are studied in structures formed by single strands, for example the hairpin formed by A₆C₆U₆. If comparisons in such cases are made with helixes formed by bimolecular associations, it is necessary to correct for initiation of the bimolecular helix first. Unfortunately, fewer data are available for loops than for fully base paired helixes. Much of the available data is discussed below. Examples of calculations of themodynamic properties for structures containing loops can be found in Xia et al.

3.1 Bulge Loops

Bulge loops are loops in which unpaired nucleotides occur on only one strand of a double helix (Fig. 8-11). In structures of natural RNA, the most common bulge contains one nucleotide, and bulges are known to be important for protein binding and tertiary folding (Peatite et al., 1981; Romaniuk et al., 1987; Flor et al., 1989; Cate et al., 1996, 1997). In DNA, bulges may be important in frameshift mutagenesis in sequences with repeating base pairs (Okada et al., 1972), and detection of bulges by gel electrophoresis can reveal mutations (Triggs-Raine and Gravel, 1990; Rommens et al., 1990). Chemical modification experiments on large RNA

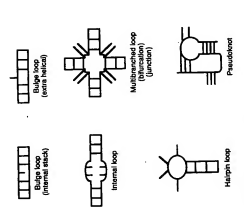


Figure 8-10 Schematic of various types of loops.

molecules (Moazed et al., 1986) suggest the structure of a bulge may be sequence dependent. Using R and Y to denote purines and pyrimidines, respectively, bulged nucleotides in N^RR sequences are usually moderately or strongly reactive. Bulged nucleotides in Y^RY sequences are usually protected from modification. NMR experiments indicate the bulged As in (GCGCAGAATTTCGCG), (GCGCAGCTCGCG), and (GCGCAAATTTGCG), are intercalated in the helix (Patel et al., 1982, 1986; Hare et al., 1986, while the bulged C in q(CAAAQAAG) dCTTTTTTG) and the bulged U in r(CUGGUGCGG) r(CCGCCAG) are extrahelical (Morden et al., 1983; van den Hoogen et al., 1988) (see Chapter 4). Thermodynamic parameters for single nucleotide bulges indicate they destabilize the helix, but with little dependence on the identity of the bulged base (Longfellow et al., 1990; Groebe and Uhlenbeck, 1989). The destabilization, however, does depend on more than the adjacent base pairs (Longfellow et al., 1990). Thus the nearest neighbor model is oversimplified for bulges. A limited number of sequences have been studied, and representative data are listed in Table 8-9.

For example, any of the middle C nucleotides in the lower strand of acrococcus could be bulged. The NMR experiments indicate such migration occurs at a rate of at least 100–1000 s⁻¹ (Woodson and Crothers, 1987).

Theoretical considerations indicate bulge loops should be more destabilizing as the number of unpaired nucleotides increases. While only a few measurements have been made on bulges larger than 1, the results are mostly in agreement with this prediction (see Table 8-9) (Longfellow et al., 1990; Yuan et al., 1979; Weeks and Crothers,

Some bulges occur in sequences that allow migration of the unpaired nucleotide.

3.2 Internal Loops

with respect to the number of loop residues on each strand. Asymmetric internal loops are less stable than symmetric internal loops (Peritz et al., 1991). A particularly important subclass of internal loop, the mismatch, contains two nucleotides. Stabilities of mismatches are important for determining stabilities of duplexes formed between Due to redundancy in the genetic code, this situation often occurs when the sequence of a probe is designed from the sequence of a gene's protein product. Modified bases, like inosine, that are less discriminating than A, C, G, and T, can be used in such cases to lessen the effect of the mismatch (Martin et al., 1985; Cheong et al., 1988; Nichols et al., 1994). Formation of large internal loops is the first step in melting of DNA (Blake and Decourt, 1987; Wada et al., 1980; Gotoh, 1983; Wartell and Benight, 1985). In RNA, internal loops of many sizes are known to occur (see Fig. 8-1). These loops allow number of nucleotides in the loop. Internal loops can be symmetrical or asymmetrical DNA hybridization probes and target sequences that are not completely complementary. 1995), and can also form binding sites (Sassanfar and Szostak, 1993; Fan et al., 1996; paired nucleotides on both strands (Fig. 8-11). The size of an internal loop is the total structural bends (Murphy and Cech, 1993) and tertiary interactions (Costa and Michel. An internal loop is formed when a double helix is interrupted by non-Watson-Crick liang et al., 1996; Dieckmann et al., 1996; Yang et al., 1996).

Considering the wide variety of possible internal loops, relatively few experimental data are available. Some measurements involve single mismatches in DNA

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37°C	
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v Increments at 37°C for Bulge Lo	
.9 Prever	ò
able 8.9	2
- u	•

Bulge Sequence	Reference	<u>∑</u> §	r _m (°C 10⁻⁴∧)	Δι _m (°C 10 ⁻⁴ Λι)	ΔΔG _{bulge, 37} (kcal mol ⁻¹)
RNA					;
poly(A.A") · polyU	Fink and Crothers (1972)	0.2-0.5			6.7
3'GCG^GCG3' 3'CGC-CGC3'	Longfellow et al. (1990)	-	8	8	3.5
s'aca ^u aca; s'coc-cacs'	Longfellow et al. (1990)	-	88	12	. 3.7
3'0CG-GCG3'	Longfellow et al. (1990)	-	39	70	3.5
s' GCG^GCGA3' 3' ACGC-CGCS'	Longfellow et al. (1990)	-	84	22	1.3
S' GCG^GUCA3' 3'ACGC-CAGS'	Longfellow et al. (1990)	-	6	2	2.0
S' ^G GGACUC ^A CG ^{AU} S' _{LA} UCUGAG·GC _A ^U	Groebe and Uhlenbeck (1989)	-	67	2	3.4
s, aggacucace ⁿ y _{ux} ucugag.ac _x u	Groebe and Uhlenbeck (1989)	-	£	9	3.5
S ^G GGACUC ^U CG ^{AU}	Groebe and Uhlenbeck (1989)	-	8	•	7.7
3'GCG ^{AA} GCG3' 3'CGC—CGC5'	Longfellow et al. (1990)	-	74	38	5.2
5'GCG***GCG3' 3'CGC—CGC5'	Longfellow et al. (1990)		1	\$	5.7
s'GCG—GCG3'	Longfellow et al. (1990)	-	38	21	3.7
DNA					
s'acafa,Gs' 3' GTT.Cs'	Morden et al. (1983)	-	<u>se</u>	4	2.6
S'ACTG^CCCATC3'		į	;	ž	. 5
3' GAC-GGGTAGS'	Woodson and Crothers (1987)	5	5	2	
3' GAC GGGTAGS'	Woodson and Crothers (1987)	0.1	37	9	2.1

The parameters Δ_{L_m} and $\Delta\Delta G_{bulge,33}$ are differences between the duplexes with and without the bulge.

1984; Aboul-ela et al., 1985; Allawi and SantaLucia, 1997, 1998). Table 8-10 lists some of these results. Substituting a mismatch for a base pair always gives a less stable duplex. In general, the destabilization is associated with a less favorable ΔH° and more favorable ΔS of duplex formation, which is consistent with the expectation of reduced oligonucleotides (Gaffney and Jones, 1989; Martin et al., 1985; Tibanyenda et al., bonding in a mismatch coupled with increased flexibility.

lizing as they increase in size (Weeks and Crothers, 1993; Peritz et al., 1991; Gralla and Crothers, 1973). For example, the melting temperatures at 10-4 M strand for Data for RNA internal loops indicate that internal loops become more destabi-

3 Double Helix Formation by Oligonucleotides with Loops

Table 8.10 Thermodynamic Parameters for Duplex Formation in 1 M NaCl by DNA Oligonucleotides Containing Mismatches'

Sequence	Mismatch	- AH° (kcal mol ⁻¹)	-4 <i>S</i> *	- A G ₁₇ (kcal mol ⁻¹)	t, (°C at 10-4M)
CA,CA,G + CT,GT,G		2.2	≘	1.7	42.9
CA, GA, G + CT, CT, C		62.8	671	2,2	40.8
CA,AA,G+CT,TT,G		0.89	8	1.2	40.1
CA, TA, G + CT, AT, G		58.6	89	6.5	36.8
$A_3GA_3G + CT_3GT_3G$	9	53.5	158	4.5	25.6
CA, TA, G + CT, GT, G	2	55.6	165	4,4	25.7
CA,GA,O+CT,AT,G	ď	52.6	<u>8</u>	4.2	23.9
CA,GA,G+CT,TT,G	5	46.7	137	4.2	22.3
CA, AA,G+CT,GT,G	ΨĊ	39.9	91	3.9	0.81
CA, AA,G + CT,AT,O	¥	36.9	101	3.7	15.0
CA,CA,G+CT,TT,G	Ե	53.2	9	3.3	1.61
CA, TA, G + CT, CT, G	5	90.0	131	3.2	5.71
CA,CA,G+CT,AT,G	ð	(40.3)	(120)	(3.1)	(6)
CA,TA,G+CT,TT,G	F	(54.6)	(167)	(2.8)	5
CA,AA,G+CT,CT,G	Ų	(35.8)	(106)	(2.9)	6
כא,כא,פ + כד,כד,פ	ဗ	(55.3)	(E)	(2.3)	(15)
S'CAACTTGATATTAATA					
3'GTTGAACTATAATTAT		102.1	289	12.4	55.8
S'CAACTTGATATTAATA					
3'GTTGAGCTATAATTAT	10	92.6	566	10.1	4.64
S'CAACTTGATATTAATA					
3'GTTGAACTATAGTTAT	10	95.5	274	10.5	50.5
S'CAACTTGATATTAATA					
3'GTTGAACTCTAATTAT	5	4.86	286	6.7	47.3
S'CAACTTGATATTAATA					
3'GTTGAATTATAATTAT	5	91.3	3	7.6	1.7
S'CAACTTGATATTAATA			٠		
3'GTTGAACCATAATTAT	¥	6:06	365	8.7	44.6
S'CAACTTGATATTAATA					
3'GTTGAACAATAATTAT	¥	92.0	267	9.26	46.2

Data for CA, XA, G + CT, YT, G sequences are from Aboul-ela et al. (1983). Data in parentheses are significantly less accurate. Other sequences are from Thanyenda et al. (1984). Similar results for d(GGTIXTITGG) + d(CCAAVAACC) have been reported by Gaffiney and Jones (1989).

(CGCA, GCG), with n = 0, 1, 2, and 3, are 58, 40, 36, and 32°C, respectively (Peritz et al., 1991). The stability of an internal loop is also dependent on the sequence in the loop (SantaLucia et al., 1991; Wu et al., 1995; Schroeder et al., 1996; Xia et al., stabilize a duplex, while other double mismatches destabilize a duplex. This sequence 1997). For example, Table 8-11 lists ΔG values for internal loops containing adjacent, identical mismatches. For this case, GU, GA, and UU double mismatches often

Chapter 8 / Conformational Changes

R N A	
Table 8.11 Free Energy Increments (ΔG_3 , in kcal mol ⁻¹) for Tandem Mismatches in RNA	
Misma	
Tandem	
-1) for	
cal mol	
37 in kc	<u>1</u>
nts (A G	1 M Na
ıcreme	Oligonucleotides in 1 M NaCl*
8.1.1 nergy Ir	nucleoti
Table 8.11 Free Energ	Oligo

		١				١						
Mismatches	↓25¢	↓98†	1 Q Å ↑	↑52†	t33†	↑88†	1251	†88†	†25†	†88†	t 52 t	↑\$\$↓
Closing bp												
S'G												
ည္ကုန္	6.4	4	-2.6	-1.3	-0.5		0.	=			6.0	<u>.</u>
의	-4.2	7	-0.7	-0.7	4.0	8.0	Ξ	4.	4.	7	5.0	1.3
S 🔄	-2.6	-0.3	0.7		Ξ		[1.9]	2.2	2.8			2.8
≲긺	-1.9	0.2	0.3		9.0		2.3				2.5	2.8
												İ

'Free energy increments are calculated as in the following example:

$$\Delta G_{j_1,\text{low}}\left(\begin{array}{c} \text{CAGG} \\ \bullet & \bullet \\ \text{GGAC} \end{array} \right) = \Delta G_{j_1}(\text{CGC}\underline{A}\underline{G}\text{GCG}) - \Delta G^*(\text{CGCGCG}) + \Delta G_{j_1}\left(\begin{array}{c} \text{CG} \\ \text{GC} \end{array} \right)$$

Most sequences had 3 bp on each side of the internal loop. Some values are averages from more than

'He et al., (1991); SantaLucia et al., (1991); Wu et al., (1995); Mathews et al., (1999). one sequence.

'Sequence with this tandern mismatch had either an unusual conformation or mixture of conformations.

dependence is thought to be due to hydrogen bonding in the stable loops (SantaLucia et al., 1991; Wu et al., 1995). The NMR studies of internal loops have revealed a variety of noncanonical interactions indicating that rules for thermodynamic stability may be complex (Wimberly et al., 1993; SantaLucia and Turner, 1993; Peterson et al., 1994;

introns (Pyle et al., 1994; Knitt et al., 1994; Strobel and Cech, 1995). Table 8-12 lists thermodynamic parameters for GU mismatches (Mathews et al., 1999). Note that the nearest neighbor, S'GU3/S'GU3', is destabilizing in most but not all contexts. This sequence dependence is a non-nearest neighbor effect. The destabilizing motif rarely nal GU pairs, and these terms are assumed to be the same as those listed for terminal The most common mismatch in RNA is GU, and often GU mismatches are conserved (Gutell et al., 1994; Michel and Westhof, 1990). This conservation may imply occurs in secondary structures of natural RNA. Additional terms are applied for termiimportance in tertiary and functional interactions, and this has been shown for group I Battiste et al., 1994; Wu and Turner, 1996; Wu et al., 1997). AU pairs in Table 8.4.

Table 8.12

Thermodynamic Parameters for Helix Propagation by G:U Pairs in RNA Oligonucleotides in 1 M NaCl*

ΔG_{j7}^{\bullet} (kcal mol ⁻¹)	-2.51	-2.11	-1.53	-1.27	-1.36	4.1.4	00.1-	-0.55	+0.30	+1.29(-1.06)*	+0.47
Δ <i>S</i> * (eu)	-32.5	-32.2	-21.9	-37.3	-24.0	-13.5	-19.3	9.8	-30.8	-51.2(-42.2)*	-44.9
ΔH° (kcal mol ⁻¹)	-12.59	-12.11	-8.33	-12.83	18.8	-5.61	66.9-	-3.21	-9.26	-14.59(-14.14)*	-13.47
Propagation Sequence	† 99 t	† & 3 †	†88±	1 5 5 1	† 5 ₹ †	1881	↓ 2 5 ↑	† & B ↑	† 53 £3 ↓	↑ <u>9</u> 8 †	†83↓

'He et al., (1991); Mathews et al. (1999). Additional terms are applied for terminal GU pairs, and these are assumed to be the same as those listed for terminal AU pairs in Table 8.4.

• GU in the contexts AGUU CGUG and UGUA has a AG", AH", and AS" of +1.29 UG

kcal mol-1, -14.59 kcal mol-1, and -51.2 cu, respectively, but in the context GGUC it has a CUGO

ΔG', ΔH', and ΔS' of -1.06 kcal mol-1, -14.14 kcal mol-1, and -42.2 eu, respectively.

3.3 Hairpin Loops

pairs (Fig. 8-11). Hairpins are widespread in structures of natural RNA (see Fig. 8-1). In to the fully base paired double helix. There is evidence that such hairpins will form Hairpin loops occur when nucleic acid strands fold back on themselves to make base DNA, there are many instances of sequences capable of forming hairpins as alternates when the DNA is placed under superhelical tension (Gellert et al., 1979; Panayotatos and Wells, 1981; Sullivan and Lilley, 1986).

namic parameters can be determined from calorimetric and spectroscopic data in a hairpin transitions are more cooperative than those of single strands, thus simplifying determination of baselines. For transitions considered two-state, ΔH° is simply derived Conformational changes involving hairpins are intramolecular. Thus thermodymanner similar to that described for single-strand stacking in Section 8.1.1. Fortunately, from (Bloomfield et al., 1974):

Unimolecular
$$\Delta H^{\circ} = 4RT_{m}^{2} \left(\frac{\partial \alpha}{\partial T}\right)_{T=T_{m}}$$
 (8-22a)

Here the partial derivative is the slope at the T_m of a plot of fraction of strands in single-strand state versus temperature. The value of $\Delta \mathcal{S}$ can then be determined from Eq. 8-7b. Equation 8-22a is a specific case of the general equation for an equilibrium with N strands (Marky and Breslauer, 1987):

N-mer
$$\Delta H^{\circ} = (2 + 2N)RT_{n}^{2} \left(\frac{\partial \alpha}{\partial T}\right)_{T-T}$$
 (8-22b)

The complete basis for the sequence dependence of hairpin stability has not yet been unraveled (Hilbers et al., 1994), although hydrogen bonding and stacking within the loop probably make contributions (Varani et al., 1991; SantaLucia et al., 1992; Serra et al., 1994). An equation that provides a reasonable prediction for the free energy increment in kilocalories per mole associated with RNA hairpin loops greater than or 1994). The free energy increment for loop formation is unfavorable, and the minimum 1995). The stability of hairpins is dependent on loop size, the sequence in the loop, and the base pair closing the loop (see Table 8-13). For example, the CG closed tetraloop More data are available for hairpins than for other loops (Table 8-13; Hilbers et al., loop size is 2 (Orbons et al., 1986, 1987; Blommers et al., 1989; Jucker and Pardi, formed by CUUCGG is unusually stable (Tuerk et al., 1988; Antao and Tinoco, 1992). equal to 4 nucleoiides is (Serra et al., 1994, 1997; Mathews et al., 1999):

$$\Delta G_{yy,loop} = \Delta G_{yy,length}^o + \Delta G_{yy,loop}^o = 0.8$$
 if first nuismatch is GA or UU (8-23)

Here $\Delta G_{\rm 31,leags}$ is the length dependence of hairpin ΔG (see Table 8-14) and $\Delta G_{\rm 31,mm}$ is the free energy increment for the first mismatch in the loop (Table 8-6). The sarcin/ricin loop from 28S ribosomal RNA has 17 nucleotides, many of which are involved in noncanonical interactions (Szewczak et al., 1993). This result suggests that the complete set of rules for hairpin stability may be complex.

3 Double Helix Formation by Oligonucleotides with Loops

Thermodynamic Parameters for Hairpin Loops **Table 8.13**

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A ₄ Senior et al. (1981a) 0.14 Na ⁴ -33.7 C ₄ Senior et al. (1981a) 0.14 Na ⁴ -40.8 C ₄ Senior et al. (1981a) 0.14 Na ⁴ -40.3 T ₄ Senior et al. (1981a) 0.14 Na ⁴ -40.3		٠,	Hilbers et al. (1985).	0.2M/Na		1360		
C ₄ Senior et al. 1988a). 0.1 M Na* -41.8 C ₄ Senior et al. 1988a). 0.1 M Na* -40.3 T ₄ Senior et al. 1988a). 0.1 M Na* -40.3 3c	CGAACG	₹	Senior et al. (1988a).	0.1M Na				
C ₄ Senior et al. (1984s), 0.1 M No46.3 T ₄ Senior et al. (1984s), 0.1 M No49.3 Gm ² C O ₂ Control (1984s), 0.1 M No49.3	3	ď	Senior et al. (1988a).	O.I.W.Na				
T ₄ Senior et al. (1988a). 0.1M Na ⁺ -49.3 Gm ³ C on comment of the senior -230		ťű	Senior et al. (1988a).	0.1M Na				
Cm ² C			Senior et al. (1988a).	0.1M No.				
	m ² C Gm ² C		1100T		280	8.73		

Table 8.14 Free Energy Increments (kcal mol⁻¹) for RNA Loops at 37°C in 1 M NaCl*

0001	Internal	Bulge	Hairpin
Size	roop	Loop	Loop
-		+3.8	
	+0.4	+2.8	1
, "		+3.2	+5.7
n 4		(+3.6)	+5.6
, v	8.1+	(+4.0)	+5.6
י ע	+2.0	(+4.4)	+5.4
, ,			+5.9
· œ			+5.6
o			+6.4

From Mathews et al. 1999. For larger loop sizes, n, use $\Delta G^*(n) = \Delta G^*(n_m) + 1.75RT \ln (n/n_m)$, where n_m is 6, 6, and 9 for internal, bulge, and hairpin loops, respectively. Parameters not derived from experimental measurements are listed in parenthleses. Farameters for sharpin loops are for from experimental measurements are listed in parenthleses. Farameters for sharpin loops are for before by GG of GG, and for loops greater than or equal to 4 assume additional stability is conferred by terminal mismatches at helix ends (see Table 8-6). A reasonable approximation for hairpin loops by terminal mismatches at helix ends (see Table 8-6). A reasonable approximation for hairpin loops per all reasonable approximation of hairpin loops penalized additionally by the minimum of 3.0 or 0.5 | NI | - N2| keal mol⁻¹. For internal loops larger penalized additionally by the minimum of 3.0 or 0.5 | NI | - N2| keal mol⁻¹. For internal loops arget penalized additionally by the minimum of 0.0 or 5 | NI | - N2| keal mol⁻¹. For each AU or GU pair clossing an internal loop, a penalty of 0.2 keal mol⁻¹ is applied in addition to the 0.45 keal mol⁻¹ penalty for terminating a helix with AU or GU. The parameter for bulge loops of one nucleotide is approximated by nearest neighbor parameters (see Table 8-4), it is assumed there is no stacking across bulges of two or more nucleotides.

'Rough average for single mismatches that are not GG and that have 2 adjacent GC pairs. The equivalent value for GG mismatches is -2 keal mol-1.

equivalent value for GG musmatones is - a real mon

Free energies of internal loops with 3 or 4 nucleotides are estimated with special rules (Xia et al.,
1999; Mathews et al., 1999).

The kinetics of hairpin formation have been studied by temperature-jump methods (Pörschke, 1974; Riesner et al., 1973; Coutts, 1971; Coutts et al., 1974; Orbons et al., 1986, 1987). Forward rates are between 10^4 and 10^3 s⁻¹ and change little with temperature. As with duplex formation, this result allows rough prediction of the rate of unfolding for a hairpin, if the equilibrium constant for hairpin formation is known or an be predicted. For example, at the melting temperature, $K = k_{\rm loc}/k_{\rm loc} = 1$. Thus the rate of unfolding of a hairpin at the $T_{\rm m}$ is also roughly $10^4 - 10^5$ s⁻¹. In a temperaturejump experiment, the ecorresponding relaxation time is given by $\tau^{-1} = k_{\rm loc} + k_{\rm ev}$, or about 5–50 μ s at the $T_{\rm m}$.

3.4 Multibranch Loops (Junctions)

Multibranch loops (junctions) occur when three or more helixes intersect (Fig. 8-11). The term loop is somewhat misleading since it is possible for these structures to exist with no unpaired nucleotides. Models for such a structure with four helixes have been studied with DNA oligonucleotides (Marky et al., 1987; Lu et al., 1992). The ΔH^{μ} for formation of one structure was simply the sum of the ΔH^{μ} values for formation of each separate helix in the structure. Thus the junction did not significantly perturb the bonding in attached helixes (Marky et al., 1987). The thermodynamic parameters for forming a four-arm junction structure from two paired duplexes are $\Delta G_{ii} = 1.1$ kcal mol⁻¹ and $\Delta H^{\mu} = 27.1$ kcal mol⁻¹ (Lu et al., 1982). A DNA junction with three helixes is stabilized by adding unpaired nucleotides to the junction (Leoniis et al., 1991). Melting of RNA involves changes in the size of a multibranch loop (Riesera and Römer, 1973; Crothers and Cole, 1978; Crothers et al., 1974).

3.5 Knotted Loops

If two nucleotides i and j are base paired, a pseudo knot is formed when a nucleotide between i and j in the primary sequence is base paired with a nucleotide not between i and j in the primary sequence is base paired with a nucleotide not between i and j (Fig. 8-11). These structures were first postulated to explain similar biochemical reactivities for IRNA and the 3' ends of certain viral RNAs from plants (Pleij et al., 1985). They have subsequently been shown to occur in oligonucleotides of appropriate sequence (Wyatt et al., 1990). It was suggested that the minimum loop sizes for the 5 and 3' loops are two and three nucleotides, respectively, for A-form RNA, and perhaps even one or two for a distorted structure (Pleij et al., 1985). Studies of oligonucleotide models are consistent with the predicted minimum loop sizes for A form, and indicate pseudoknot stability is dependent on loop size and sequence (Wyatt et al., 1990). The presence of Mg²⁺ or high Na⁺ concentrations also stabilizes pseudoknots (Wyatt et al., 1900).

3.6 Large Loops

Loops larger than 12 nucleotides are rarely seen in nucleic acid structures. Nevertheless, the properties of large loops are important for successful predictions of nucleic acid structure, and for predicting the melting behavior of nucleic acids. Unfortunately, it is difficult to experimentally determine the properties of large loops, especially given the number of possible permutations. Theoretical considerations, however, provide a reasonable approximation for the ΔG for initiation of large loops (Jacobson and Stockmayer, 1950). Considerations of conformational entropy effects lead to the following equation for the ΔG° of initiation of a large loop with N nucleotides:

$$\Delta G_{\text{loop}}^{\circ}(N) = \Delta G_{\text{loop}}^{\circ}(n) + aRT \ln(N/n)$$
 (8-24)

Here $\Delta G_{\text{loop}}^{*}(n)$ is the experimentally determined ΔG^{*} for a loop of n nucleotides, and a is a constant. Theoretical considerations suggest the value of a is about 1.75 (Fisher, 1966). Experiments on bulge and internal loops are consistent with Eq. 8-24

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(Weeks and Crothers, 1993). A set of loop parameters derived from experimental and theoretical results is given in Table 8-14.

Reactions involving covalently closed circular nucleic acids provide another situation where the free energies of large loops are probably important. For example, the melting temperature of a covalently closed circular DNA of 26 nucleotides that forms a dumbbell-shaped structure is more than 30°C higher than for the same sequence with a single break in the sugar-phosphate backbone (Eric et al., 1989). This difference corresponds to a more favorable ΔG_3^+ , of 7 kcal mol⁻¹ for folding into the dumbbell. A large fraction of this favorable free energy is expected from the conformational constraints placed on a melted loop when it is covalently closed (Jaeger et al., 1990). The same effect can also largely account for the enhanced binding observed between oligomers when one is a covalently closed circle (Prakash and Kool, 1992). While such a conformational effect might be expected to provide a more favorable entropy for folding, a more favorable enthalpy is actually observed (Eric et al., 1989; Prakash and Kool, 1992). This apparent anomaly is seen even in reactions of small molecules, however (Jencks, 1975).

4. DOUBLE HELIX FORMATION WITH POLYNUCLEOTIDES

An understanding of helix formation in oligonucleotides provides a basis for understanding the properties of nucleic acid polymers. It is likely that interactions important in oligomers will be major factors determining polynucleotide properties, but that new interactions will also be important. Tertiary interactions are expected to be weaker interactions, and there is some experimental evidence to support this (Crothers et al., 1974; Banerjee et al., 1993; Jaeger et al., 1993; Laing and Draper, 1994). The most extensively studied cases of melting of nucleic acid polymers are tRNA and DNA. In both cases, melting typically starts with formation of large loops.

4.1 Transfer RNA

The experimentally determined pathway for melting of Escherichia coli formylmethionine tRNA in 0.17 M Na⁺ is shown in Figure 8-12 (Crothers et al., 1974). First, tertiary interactions and the dihydrouridine stem melt, followed by the T^UC stem, the anticodon stem, and finally the acceptor stem. The order of melting for other tRNAs may be different, indicating a sequence dependence (Riesner and Römer, 1973).

The kinetics of melting for RNA are similar to expectations from oligomers. The kinetics of melting for RNA are similar to expectations from oligomers (Riesner and Melting for the Ray). The $T\Psi C$ and anticodon hairpins typically melt with relaxation times of $10-100~\mu s$. Melting of the acceptor stern is associated with a relaxation time of about 1 ms, presumably because it is the final stem to melt and therefore opens a large hairpin loop. Melting of the least stable helix, the dihydrouridine stem, is associated with relaxation times of about 1-10~m s. This time dihydrouridine stem, is associated with relaxation times of about 1-10~m s. This time

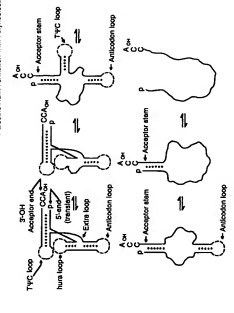


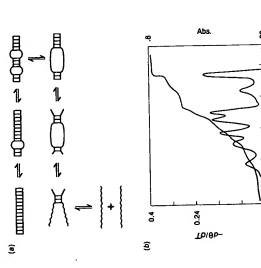
Figure 8-11
Pathway for melting of *E. coli* formylmethionine tRNA in 0.17 *M* Nu* [Reprinted Pathway for melting of *E. coli* formylmethionine taylor. W., and Shulman, with permission from Crothers, D. M., Cole, P. E., Hilbers, C. W., and Shulman, R. G., The Molecular Mechanism of Thermal Unfolding of *Escherichia coli* Formylmethionine transfer RNA. *J. Mol. Biol.* 87, 63-68, Copyright ©1974. by permission of the publisher Academic Press Limited London.] The dihydrounidine helix and tertiary interactions melt first, followed in succession by the *TVC*, anticodon, and acceptor stem helixes.

is much longer than expected from model hairpins, and is associated with concomitant disruption of tertiary structure. This long relaxation time is one example of effects in polymers that are not expected from simple model systems.

4.2 DNA

A typical differentiated melting curve and pathway for melting of DNA are shown in Figure 8-13 (Blake and Decourt, 1987; Wada et al., 1980; Gotoh, 1983; Wartell and Benight, 1985; Lerman et al., 1984). Depending on sequence and length, DNA can begin melting from the ends or the middle. As temperature is raised, AT rich regions melt first, leaving internal loops in long DNA. Typically, loops with nucleotides from 100–350 pare formed. A single large internal loop of n nucleotides is more favorable than two separate internal loops containing a total of n nucleotides (see Table 8-14). Thus loops coalesce. Occasionally, it is necessary to also consider intermediates with hairpin loops (Wartell and Benight, 1985). The result, as shown in Figure 8-13,

1987). For sequences longer than about 300 bp, separation of strands can be neglected melting behavior for any DNA sequence. Alternative sets of parameters have also been suggested (Wada et al., 1980; Gotoh, 1983; Wartell and Benight, 1985; Breslauer et Wetmur, 1989; Doktycz et al., 1992). Parameters for loops have been determined by in these predictions, but it must be included for shorter sequences (Wartell and Benight, is a series of transitions as the temperature is raised. The parameters in Table 8-4 coupled with parameters for internal and hairpin loops, should be able to predict the al., 1986; Lerman et al. 1984; Klump, 1990; Delcourt and Blake, 1991; Quartin and measuring the melting of DNAs that have AT rich regions inserted (Blake and Delcourt,



temperature and differentiated melting curves for the 1630 bp Hinf I DNA Molecules: A Comparison of Theory with Experiment, Phys. Rep., 126, 67–107. Copyright ©1985 with permission of Elsevier Science-NL, Amsterdam, The Netherlands.] restriction endonuclease fragment of plasmid pBR322. [Reprinted from Wartell, R. M. and Benight, A. S., Thermal Denaturation of (a) Typical pathway for melting of DNA. (b) Absorbance versus Figure 8-12

Double Helix Formation with Polynucleotides

1985). In several cases, predictions are reasonably successful (Wartell and Benight,

1985; Steger, 1994).

DNAs of quasirandom sequence with $0.3 < F_{cc} < 0.7$ and $0.02 \le [Na^+] \le 0.4M$ is simple equations have been developed empirically to approximately predict 1, from the fraction of GC content, $F_{\rm GC}$ (Marmur and Doty, 1962). One that is useful for long For long DNA sequences with relatively random distributions of nearest neighbors, (Blake, 1996)

$$I_{\rm m}^{\rm o}(C) = 193.67 - (3.09 - F_{\rm GC})(34.64 - 6.52\log[Na^+]).$$
 (8-25)

Another that includes the effect of duplex length, D, and the percentage of mismatches, P, but neglects the effect of GC content on the salt dependence is (Wetmur, 1991)

$$I_{\rm m}(^{\circ}{\rm C}) = 81.5 + 41F_{\rm GC} + 16.6\log_{10}\left(\frac{[Na^{+}]}{1.0 + 0.7[Na^{+}]}\right) - \frac{500}{D} - P$$
 (8.26)

This equation is valid up to 1 M Na+. The analogous equation for RNA is

$$t_{\rm m}(^{\circ}{\rm C}) = 78 + 70F_{\rm GC} + 16.6\log_{10}\left(\frac{1{\rm Na}^{4}!}{1.0 + 0.7!{\rm Na}^{4}!}\right) - \frac{500}{D} - P$$
 (8-27)

An interesting application of DNA melting is the use of gels containing denaturant gradients to separate DNAs of similar length but different sequence, and to detect single base pair changes between DNAs (Lerman et al., 1984). These applications arise from a large decrease in gel mobility when denaturation induces an internal loop. Predictions for comparison with experiment are possible since denaturants such as urea and formamide mimic a temperature increase (Lerman et al., 1984; Klump and Burkart, 1977).

that of oligomers (Wetmur and Davidson, 1968; Bloomfield et al., 1974; Cantor and The kinetics of association and dissociation for large DNAs is quite different from Schimmel, 1980; Studier, 1969; Record and Zimm, 1972). This results from sequence complexity and large size. For example, association rates are affected by intramolecular helix formation and by incorrect intermolecular helix formation. Thus, when two separated strands of a large DNA are quickly cooled, they are kinetically trapped in nonnative structures and essentially never able to reform the perfectly matched helix. k, for this association has been found experimentally to be given by (Wetmur and Under conditions where the perfectly matched helix can be formed, the rate constant Davidson, 1968; Wetmur, 1991)

$$k_2 = \frac{k_u \sqrt{L}}{N} \tag{8-28}$$

Here k_{μ}' is the nucleation rate constant, L is the length of the shortest strand participating in duplex formation, and N is the complexity of the sequence. Complexity has been defined as "the total number of base pairs present in nonrepeating sequences" (Wetmur, 1991) or "the length of DNA needed to contain one copy of the entire sequence" (Cantor

1991). The kinetics of dissociation near the $T_{\rm m}$ for large DNAs occurs in the 100–1000 s time range and cannot be fit with a simple functional form. This observation is thought to be due to large frictional effects in the unwinding process and is avoided in vivo by the actions of nicking-closing enzymes (topoisomerases) and gyrases that break and Schimmel, 1980). The rate constant k', is a complicated function of temperature and other conditions, approaching zero at the T_m of a large DNA. It is maximal roughly 25°C below the $T_{\rm m}$, where a typical value is $3.5\times10^3\,M^{-1}\,{\rm s}^{-1}$ in 1 M NaCl (Wetmur, phosphodiester bonds to allow easier rotation.

4.3 Nucleic Acid Hybridization

arrays on solid support can detect thousands of sequences simultaneously (Schera et al., 1994). The effects of solid supports, however, have not been studied in detail (Wetmur, detecting specific sequences in complex mixtures. For example, the polymerase chain reaction (PCR) and various blotting techniques rely on specific binding of short or long nucleic acid primers or probes to target sequences (Wetmur, 1991). Oligonucleotide factors, including the $T_{\rm m}$ and kinetics for the binding and for the disruption of structure in individual sequences. The principles described above can often be used to predict optimum conditions for experiments requiring hybridization (Wetmur, 1991; Steger, Nucleic acid hybridization and methods that rely on it are the methods of choice for 1995; Chee et al., 1996). The best conditions for specificity are dependent on many

activation energy for probe dissociation, $E_{\Lambda_{MR}}$, the melting temperature T_m (in K) for ter that is important for a given experiment. For example, PCR relies on dissociation of be a little, typically 10° C, below $T_{\rm n}$. For probing blots, the kinetics of dissociation is the important parameter, since unhybridized probe is removed by washing for a given ciation temperature T_d as the temperature at which one-half of the correctly matched hybrid is released in a time t_{wat} . The parameter T_a can be predicted from the expected the hybrid duplex at the probe concentration, and the half-life for probe dissociation at In applying the principles described above, it is important to recognize the paramehelixes, so the system should be brought to a temperature above T_m to insure complete dissociation. For the annealing step, specificity is important, so the temperature should time period. For this application, Wetmur (Wetmur, 1991) suggests defining a disso-

$$\ln(t_{wath}/t_{1/2}) = (E_{A,off}/R)(T_d^{-1} - T_m^{-1})$$
 (8-29)

1.4 Tertiary Interactions

The term "tertiary interaction" is used in several different ways. Sometimes it refers to noncanonical interactions, for example a hydrogen bond to a phosphate group as sometimes seen in hairpin and other loops (Varani et al., 1991; Heus and Pardi, 1991; SantaLucia et al., 1992; Pley et al., 1994). Here we use a more restricted definition interaction is any interaction, direct or indirect, between a nucleotide k between i (Chastain and Tinoco, 1991). If two nucleotides i and j are paired, then a tertiary

and j in the sequence, and another nucleotide & which is not between i and j. Thus a pseudoknot (Fig. 8-11) is one form of tertiary interaction. Another is metal ion mediated interactions between nucleotides k and l (Lu and Draper, 1994). One approach to understanding tertiary interactions is to study binding of oligonu-1988, 1989b; Herschlag and Cech, 1990; Bevilacqua et al., 1992), fluorescence Bevilacqua et al., 1992; Sugimoto et al., 1989a), equilibrium dialysis (Bevilacqua and Turner, 1991), and gel retardation (Pyle et al., 1990). The last method is the fastest, but is not completely understood. When using gel retardation, it is important to allow sufficient incubation time before loading samples on the gel (Bevilacqua and Turner, 1991; Pyle et al., 1994). Another method for studying tertiary interactions is to monitor the effects of site directed mutation on RNA folding and function (Murphy and Cech, cleotides to natural RNAs. The methods employed include kinetics (Sugimoto et al., 1994; Costa and Michel, 1995). Little is known about the thermodynamics of tertiary interactions. One interaction that has been identified involves hydrogen bonding to 2'-0H groups for recognition of an oligomer substrate by a group I ribozyme (Sugimoto et al., 1989b; Pyle and Cech, 1991; Bevilaqua and Turner, 1991; Strobel and Cech, 1993). Each such interaction can provide a free energy increment of about 1 kcal mol-1 (Bevilacqua and Turner, 1991). In one case, the interaction involves a hydrogen bond from the H of the 2'-OH to the NI of an A (Pyle et al., 1992).

to put strain on the phosphodiester bond between G and U. This bond is the site of cleavage in the second step of group I splicing, and such strain may provide a catalytic advantage (Moran et al., 1993). Such substrate destabilization has also been reported in Unfavorable tertiary interactions are also possible. For example, UCdGU binds to a group I ribozyme at least 30-fold more weakly than UCdG (Moran et al., 1993). A model of the binding site (Michel and Westhof, 1990) suggests that it is designed a ribozyme model system for the first step of splicing (Bevilacqua et al., 1994; Narlikar et al., 1995). Favorable and unfavorable tertiary interactions are presumably sensitive to the overall folding of an RNA. Thus they can give rise to cooperativity and anticooperativity in binding of substrates. Such effects have also been observed with a group I ribozyme Bevilacqua et al., 1993; McConnell et al., 1993).

5. ENVIRONMENTAL EFFECTS ON HELIX STABILITY

5.1 Salt Concentration

stability. Up to about 0.2 M Na+, the increase in Tm is linear with log [Na+] (see Eq. 11found $dT_m/d \log[Na^+] = 18.30 - 7.04F_{oc}$ (F_{cc} is the fractional GC content). From studies of subtransitions in the melting of lambda DNA, Blake and Haydock (1979) A theoretical treatment of salt effects on helix stability for polyelectrolytes is given in Chapter 11. Here we focus on experimental results. In solutions containing only Na tor 20). The rate of increase depends on base composition. For example, from fitting data obtained on different, natural DNAs (Owen et al., 1969), Frank-Kamenetskii (1971) similar ions, increasing salt concentration up to about 1 M continuously increases helix

5 Environmental Effects on Helix Stability

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proposed a similar equation: $dT_{\rm m}/d\log[{\rm Na}^+]=19.96-6.65F_{\rm GC}$. Thus, the $T_{\rm m}$ of a deoxypolynucleotide with only AT base pairs increases almost $20^{\circ}{\rm C}$ for every factor of 10 increase in [Na+].

1962). This order correlates with the effect of these ions on the solubility of the bases Robinson and Grant, 1966). The better denaturants are most effective in enhancing At salt concentrations above 1 M, addition of salt lowers the T_m of DNA. The lowering is relatively independent of cation, but is strongly dependent on anion with CCI, COO - > SCN - > CIO7 > CH, COO - > Br -, CI - (Hamaguchi and Geiduschek, base solubility.

addition of Mg^{2+} initially increases T_m . The effect saturates at about $10^{-3}-10^{-2}M$, however. Thereafter, addition of Mg^{2+} lowers T_m (Blagoi et al., 1978). Somewhat counterintuitively, addition of Nat in the presence of enough Mg2t to neutralize the backbone charges results in a lowering of $T_{\rm m}$. This lowering of $T_{\rm m}$ is expected from concentrations of Mg2+, the effective charge on single strands is larger than on double strands due to counterion condensation. Thus, single strands are stabilized more by theoretical considerations (Record, 1975; Manning, 1978). In particular, at saturating There are few studies of the effect of Mg2+ concentration on polynucleotide melting (Riesner and Römer, 1973; Blagoi et al., 1978; Krakauer, 1974). As with Na+ increases in ionic strength because of Debye-Hückel screening effects.

ixes. Qualitatively, the effects are similar to those observed for polynucleotides. For example, dT_m/d log [Na+] for dGCATGC and dGGAATTCC are 11 and 12°C, respectively (Williams et al., 1989; Erie et al., 1987), only a little less than predicted for polymers with the same GC content. This result is somewhat surprising since theoretical considerations indicate less charge will be neutralized in oligonucleotides than in polynucleotides because of the reduced charge density at the ends (Record and Lohman, One interesting question is: At what length does an oligomer behave much like a polymer? This question has not been investigated in detail. The values of $dT_{\rm m}/d\log[{
m Na}^{-1}]$ for A, U, and poly (A) poly (U) are similar, however: 17.4 and 19.6°C, respectively 1978; Olmsted et al., 1989). Relatively little experimental data is available, however. Relatively little is known about salt effects on stabilities of oligonucleotide he-(Hickey and Turner, 1985b).

5.2 Solvent Effects

the concentration of cosolvent required to give 50% denaturation of T4 DNA at 73°C (Levine et al., 1963) and by measuring the effect of cosolvents on thermal denaturation and the effect on Tn.. This finding may reflect the very different conditions and assays (Lerman et al., 1984). Typically, the T_m of a double helix will be a linear function of cosolvent concentration (Klump and Burkart, 1977; Hickey and Turner, 1985b). For example, the $T_{
m m}$ values of calf thymus and salmon sperm DNA decrease 2.5° C M^{-1} urea (Klump and Burkart, 1977). The effects of cosolvents have been studied by measuring curves of various oligonucleotides (Hickey and Turner, 1985b; Albergo and Turner, 1981). Some of these results are listed in Table 8-15. Somewhat surprisingly, there is no simple correlation between the concentration required for 50% denaturation at 73°C Addition of cosolvents to aqueous solutions of nucleic acids typically destabilizes the ordered form, and this is often used as a substitute for denaturation by temperature

Effects of Denaturants on Duplex Stability Table 8.15

	of T4 DNA	Molarity for 50% denaturation of T4 DNA at 73°C, 43 mM	Δ¢,(°C)	Δζ _n (°C) at 10 mol % for 18.7M Olien in 1.M NaCl
		0		
Cosolvent	Predicted"	Observed	۸,۵,۲	(dGC),
Aicohols				
Methanol	3.9	3.5		9'9
Ethanol	7	1.2	8.3	Ξ
1-Propanol	0.47	0.54	9.1	8.4
2-Propanol	20.0	06'0	8.4	
Ethyleneglycol	1.7	2.2	7.7	
Glycerol		<u>~</u> :	7.9	
Cyclohexyl alcohol		0.22		
Phenoi		80:0		
Other Compounds				
Pyridine		60'0		
1,4 Dioxane		29.0	18.7	
Formamide	<u>.</u>	6:1	14.8	12.0
N, N dimethylformamide	0.54	09'0	16.9	~12
Urea	Ξ	0.1	17.8	13
Acetonitrile		1.2		
TritonX-100		2 601 ^		

'Herskovits and Hartington, (1972); Herskovits and Bowen, (1974).

Levine et al., (1963).

Hickey and Turner, (1985).

'Albergo and Turner, (1981).

used in the experiments. For example, the cosolvent required for 50% denaturation was was measured at 1 M Na with optical melting curves. The cosolvent concentrations required for 50% denaturation correlate well with enhancement of base solubility and this has predictive value (see Table 8-15). This correlation seems reasonable since the bases are more exposed to solvent in the denatured form, and favorable interactions (Levine et al., 1963; Herskovits and Harrington, 1972; Herskovits and Bowen, 1974), measured for an ionic strength of 0.04 with an antibody assay, whereas the effect on $T_{
m c}$ between bases and cosolvent favor denaturation.

5.3 pH

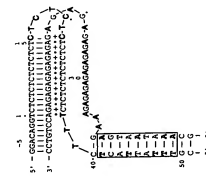
At lower and higher pH values, stability is decreased (Bloomfield et al., 1974; Blake, 1996). At low pH, bases in the single strand bind more protons than in the duplex, thus precluding normal hydrogen bonding and increasing charge repulsion. At both The stability of most double helixes is relatively insensitive to pH between 5 and 9. thus favoring single strand. At high pH, guanine, thymine, and uracil are deprotonated,

This results from hydrogen bonding schemes in which a proton is shared between two pairs are hard to form between stretches of AT pairs. In RNA, however, C. C. pairs have bases, for example A · A + . The protonated pairs do not have to be adjacent. Poly(dCT) 1980). Duplex formation by C·C* pairing has also been observed at the ends, but not the middle, of mixed deoxy sequences. For example, d(C,A,T,C,) forms such been observed in the middle of the duplex (CGCCCGCG), (SantaLucia et al., 1991). Inple-strand helixes involving C·G·C⁺ pairing have also been observed to form upon protonation of C (Lee et al., 1979). Little is known about the pH dependence of structure for other sequences, but there are suggestions that interesting effects will be discovered (Topping et al., 1988; Kao and Crothers, 1980; Legault and Pardi, 1994). For example, it has been shown that 5S rRNA undergoes a tertiary structural switch between pH 7 and 8 depending on conditions of salt and temperature (Inman, 1964; Hartman and Rich. 1965; Holcomb and Timascheff, 1968; Adler et al., 1969; Guschlbauer, 1975). forms a duplex with C·C⁺ pairs alternating with unpaired thymines (Gray et al., duplexes (Gray et al., 1984), but $d(A_{10}C_4T_{10})$, $d(AACC)_s+d(CCTT)_s$, and $d(A_sC_sA_{e})$ Some sequences form double helixes only at low pH. For example, poly(A), poly(dA), poly(C), and poly(dC) form duplexes with transition midpoints between pH + d(T,C,T,) do not (Edwards et al., 1988). This context dependence indicates C.C. and 8 (Kao and Crothers, 1980).

6. TRIPLE HELIXES

overall structure is referred to as H-DNA. It has also been proposed that pairing to et al., 1988; Praseuth et al., 1988). These ideas may provide the basis for therapeutics Hydrogen bonded base triples are found in tRNA (see Fig. 4.13). A triple-stranded structure consisting of T.AT and C. GC base triples has been proposed for regions 1987; Voloshin et al., 1988; Htun and Dahlberg, 1988; Johnston, 1988; Wells et al., ,988; Mirkin and Frank-Kamenetskii, 1994). When this triple strand forms, the remaining (dA-dG), strand is released from pairing as shown in Figure 8-14, and the give triple strands could regulate transcription (Cooney et al., 1988; Miller and Sobell, of DNA containing (dT-dC), '(dA-dG), repeats (Christophe et al., 1985; Mirkin et al., 1966) and be used for site specific modification or cleavage of nucleic acid (Strobel that bind as a third strand.

Only sequences that are largely polypurine and polypyrimidine have been observed 1957; Stevens and Felsenfeld, 1964), A·UA (Broitman et al., 1987), T·AT (Riley U·CG (Michel et al., 1990; Michel and Westhof, 1990), G·TA (Yoon et al., 1992; Griffin and Dervan, 1989), G·CG (Lipsett, 1964; Lipsett, 1963), and A·GC (Chastain and Tinoco, 1992). Triple-strand helixes have also been observed with 2'-5' linked oligonucleotides (Jin et al., 1993). The most thoroughly studied triplex is poly(A) · 2 Michelson et al., 1967). Known triple-strand pairings include: U · AU (Felsenfeld et al., to form triple strands (Lee et al., 1979; Wells et al., 1988; Felsenfeld and Miles, 1967; et al., 1966), C+GC (Lee et al., 1979; Lipsett, 1964), T·CG (Yoon et al., 1992).



Science 241, 1791-1796. Copyright Htun, H. and Dahlberg, J. E. (1988) break to allow formation of a triple structure, Watson-Crick base pairs Schematic of structure for H-DNA Advancement of Science.] In this Reprinted with permission from ©American Association for the Figure 8-13

The stability of the triplex relative to duplex and single strands depends on temperature and salt concentration as shown in the phase diagram in Figure 8-15 (Stevens and Felsenfeld, 1964; Krakauer and Sturtevant, 1968). As expected, high-salt favors triplex which is true for Mg2+ as well as for Na+ (Felsenfeld et al., 1957). For example, in from optical mixing curves (Felsenfeld et al., 1957; Stevens and Felsenfeld, 1964). region 2 in Figure 8-15 the combination of poly(A) · 2 poly(U) + poly(A) is more stable than 2 [poly(A) poly(U)]. Thus any duplexes present will disproportionate to 1965; Krakauer and Sturtevant, 1968). The stoichiometry of this complex was obtained poly(U) (Felsenfeld et al., 1957; Stevens and Felsenfeld, 1964; Ross and Scruggs, form triplex and single-strand poly(A):

$2[poly(A)poly(U)] \rightleftharpoons [poly(A)2 poly(U)] + poly(A)$

 ΔH° for formation of triplex from duplex and poly(U) is about -4 kcal mol⁻¹ (Ross and Scruggs, 1965; Krakauer and Sturtevant, 1968). The time required for triple helix The ΔH° for this reaction is about 4 kcal mol $^{-1}$ (Krakaucr and Sturtevant, 1968). The formation ranges from less than 1 to many minutes depending on length and conditions (Felsenfeld et al., 1957; Pörschke and Eigen, 1971).

7. G-QUARTETS

Hydrogen-bonded G quartets are known to occur in the telomeric regions at the ends of chromosomes (Williamson et al., 1989), and G tetraplexes can form from G monomers (Gellert et al., 1962; Pinnavaia et al., 1978), G rich DNA and RNA oligomers (Sen and Gilbert, 1988; Sundquist and Klug, 1989; Sen and Gilbert, 1990; Kim et al., 1991;

Dependence of f, on the concentration of Na* at neutral pH in the absence of divalent ions for Reprinted by permission of John Wiley & Sons, Inc.) The lines correspond to the following reactions: (1) poly($A \cdot U$) \rightarrow poly(A) + poly(U), (2) poly($A \cdot U$) \rightarrow I/2 poly($U \cdot A \cdot U$) + I/2 poly($U \cdot A \cdot U$) \rightarrow poly($U \cdot A \cdot U$) \rightarrow poly($A \cdot V$) + poly(V), (4) poly($U \cdot A \cdot U$) \rightarrow poly($A \cdot V$) Complexes, Krakauer, H. and Sturtevant, J. M. Biopolymers, 6. 491-512. Copyright @1968. various reactions of polyA and polyU [Heats of Helix-Coil Transactions of PolyA-PolyU coly(U). Note that the topmost region corresponds to single stranded poly(A) + poly(U). Figure 8-14

Hardin et al., 1991; Jin et al., 1992; Lu et al., 1993), and polymers (Zimmerman et al., 1975). Formation of the G tetrad is favored by K+ relative to Na+ (Gellert et al., 1962; Pinnavaia et al., 1978; Sen and Gilbert, 1990; Hardin et al., 1991; Hud et al., 1996). It has been suggested that formation of the G tetrad may be important for several biological processes (Williamson et al., 1989; Sen and Gilbert, 1988; Sundquist and Klug, 1989; Sen and Gilbert, 1990; Williamson, 1994). The thermodynamics of tetraplex formation has been studied for several oligomer sequences (Jin et al., 1992; Lu et al., 1993). In the presence of K^+ , ΔG_2^* , and ΔH^o per tetrad appear to be -2 to -3 kcal mol-1 and -20 to -30 kcal mol-1, respectively (Jin et al., 1992).

8. PREDICTING SECONDARY STRUCTURE

the methods are also applicable to DNA. One goal of these methods is to predict the base pairing, or secondary structure, for single-stranded chains. Experiments indicate A major application of our knowledge of conformational changes is the prediction of nucleic acid structure from sequence. Most applications have involved RNA, but that tertiary structure is less stable than most secondary structure (Crothers et al.,

8 Predicting Secondary Structure

1984; Tinoco et al., 1971). The structure predicted to be most prevalent at equilibrium 1974; Bancrjee et al., 1993; Jaeger et al., 1993; Laing and Draper, 1994). Thus as a that summing the free energy changes for such interactions will provide an approximation for the stability of a given structure (Turner et al., 1988; Papanicolaou et al., however, that the relative concentrations predicted for various species is quite sensitive to the ΔG values in the calculation. The predicted equilibrium constant between two species at 37°C changes by a factor of 10 for every 1.4 kcal mol $^{-1}$ in ΔG° . Thus current termined by strong, local interactions such as stacking and hydrogen bonding suggests is then the one with the lowest free energy (see Eqs. 8-3 and 8-7a). It should be realized, first approximation, tertiary interactions can be neglected when predicting secondary structure. The conclusion that stabilities of small nucleic acid structures are largely depredictions should be considered rough approximations.

predicted well with a nearest neighbor model. Thus, this approximation provides a reasonable treatment for helical regions. Much less is known about the sequence As discussed in Section 2, stabilities of oligonucleotides without loops can be dependence of stability for structures with loops, and current methods largely neglect data become available. Even with these limitations, the loop parameters listed in Table this sequence dependence. This restriction can be eliminated when more experimental 8-14 are useful for predicting RNA secondary structure.

from sequence. The reason is that the time required to try every possibility is usually randomly, the number of valid structures goes approximately as 1.8" (Zuker and an 80 nucleotide sequence. To circumvent this problem, clever computer algorithms and Stiegler, 1981; Williams and Tinoco, 1986; Nussinov et al., 1982; Zuker, 1989; Mathews et al., 1999; Rivas and Eddy, 1999). These algorithms, however, require If sequence dependent free energy parameters were available for every possible loop, it would still be difficult to completely predict the most stable conformation enormous. For example, for a sequence of N nucleotides with A, C, G, and U occurring Sankoff, 1984). If ΔG could be calculated for 1000 structures every second, it would take about 1010 years, roughly the age of the universe to try all valid possibilities for have been written that avoid trying every possibility (Papanicolaou et al., 1984; Zuker additional approximations. Two of the approaches are discussed below.

8.1 Combinatorial Algorithms

to be low in free energy. Nevertheless, the time required is large because the number Combinatorial algorithms develop a list of all helixes that can be formed from a sequence (Papanicolaou et al., 1984; Gouy, 1986). These algorithms can include knotted structures. The algorithms then try combinations of these helixes in search of the lowest free energy. Various tricks are used to avoid computing combinations that are not likely of helixes, L, and the number of combinations grow approximately as N2 and 2L, respectively.

8.2 Recursive Algorithms

Recursive (or dynamic) algorithms usually make the approximation that if two nucleotides pair, then any nucleotide between them in the primary sequence can only pair

Chapter 8 / Conformational Changes

with other nucleotides between the originally paired nucleotides (Zuker and Sankoff, 1984; Williams and Tinoco, 1986; Nussinov et al., 1982). Thus knots are usually not allowed, but the other loops shown in Figure 8-11 are allowed. This approximation permits the lowest free energy structure to be found from consideration of the lowest free energy structure for each possible subfragment of the sequence. The computation for each new subfragment makes use of the computations for each smaller subfragment, which makes the algorithms recursive, and therefore fast. The time required typically grows as N^3 or N^4 , depending on the generality of the free energy parameters. When knots are included in a recursive algorithm, the time grows as N^6 (Rivas and Eddy,

The most popular recursive algorithm can use more than thermodynamic parameters for deducing structure (Zuker and Stiegler, 1981; Zuker, 1989; Mathews et al., 1999). For example, if it is known that a given nucleotide is susceptible to nuclease cleavage, the predicted structure can be forced to include this. Thus experimental data can play a role in the process.

8.3 Suboptimal Structures

Except for short sequences, free energy minimization may never lead reliably to the exact secondary structure, because many approximations are mandated by experimental and computational considerations. Even if it were possible to predict the most stable structure, it is likely that other structures will be important for understanding function. For these reasons, both combinatorial and recursive algorithms have been designed to permit prediction of suboptimal structures (Steger et al., 1984; Williams and Tinoco, 1986; Zuker, 1989; Gouy, 1986). The output from these programs can then be tested against experimental data (Mathews et al., 1997) or used to design experiments aimed at testing various possibilities. If more than one sequence is known for molecules with similar functions, then comparison of suboptimal structures may help identify the true structure and features required for function (Lick et al., 1996; Mathews et al., 1997). Figure 8-16 provides an example of a sequence with two structures that are similar in free energy. Programs for folding RNA are now available on the WEB (e.g., http://ma.chem.rochester.edu, and http://www.ibc.wustl.edu/~zuker).

8.4 Evaluating Predictions

The performance of free energy minimization procedures can be tested by comparing predicted structures for various RNA sequences with those deduced from sequence comparisons. In tests of a recursive algorithm on various RNA sequences, roughly 70% of the known base pairs are present in the free energy minimized structure (Mathews et al., 1999). The best computer-generated suboptimal structure has roughly 85% of the known base pairs (Mathews et al., 1999). Considering the lack of experimental data for many free energy parameters, the required approximations, and the difficulty of the problem, the agreement between known and energy minimized structures is surprisingly good. This finding supports the assumption that secondary structure is largely determined by local interactions and is not very dependent on tertiary structure.

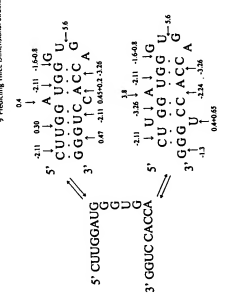


Figure 6-13

Faurple of an RNA sequence with two secondary structures of similar free energy.

The sequence is a subfragment of SS rRNA from Philosamia cynthia ricini. A program that gives suboptimal structures will predict both structures.

Presumably, performance will continue to improve as more parameters are measured

experimentally.

It may also be possible to improve predictions of structure from sequence by It may also be possible to improve predictions. From the forward and reverse rate adding considerations from the kinetics of folding. From the forward and reverse rate constants of hairpin formation, it is clear that a nucleic acid is similar to a computer algorithm in that it does not have time to try all possible pairings. Thus the pathway of folding must also be coded into the sequence. So far, only the folding of tRNA has been studied in detail (Riesner and Römer, 1973; Crothers and Cole, 1978; Crothers et al., 1974). Therefore general rules for folding are not clear yet. Preliminary attempts have been made, however, to introduce kinetics into algorithms for predicting secondary structure (Gultyaev et al., 1995; Schmitz and Steger, 1996).

9. PREDICTING THREE DIMENSIONAL STRUCTURE

Once the secondary structure of a nucleic acid is known, the next challenge is to predict the 3D structure. Good models are available for the 3D structures of double helixes (see Chapter 4), and determination of all the double helical regions greatly constrains possible 3D foldings. It is still necessary, however, to determine folding between helical regions. Presumably, this will be determined by the same factors that are important for

energy changes measured for stacking in oligonucleotides. The first correlation involves planar. This lack of planarity should lead to even less tendency to stack. Inspection of Since the factors governing 3D and secondary structure seem similar, it should be possible to use information from studies of conformational changes in oligonucleotides and polynucleotides to help predict 3D structure. The structure of yeast phenylalanine IRNA shown in Figure 8-17 illustrates three correlations of 3D structure with free single-strand stacking. Table 8-1 indicates that single-strand AA and CC sequences have considerable stacking. In the tRNA crystal structure, the sequences 35AA36 and 74CC75 are both stacked. Conversely, Table 8-1 indicates single-strand UU sequences do not stack. Dihydrouracil, D, is similar to uracil, except the pyrimidine ring is not Figure 8-17 shows that both dihydrouracils in the sequence 16DD17 are unstacked.

type of interaction is also associated with a large favorable free energy change in an The second correlation involves coaxial stacking of adjacent helixes. In the tRNA crystal structure, the base pairs 7UA66 and 49CG65 are stacked on each other. This oligonucleotide model system (Walter et al., 1994; Walter and Turner, 1994)

sequences are favored at positions where the backbone turns. While limited, the results is a turn in the sugar-phosphate backbone. These are the only places in the structure where sharp turns occur at the ends of helical regions, which suggests weakly stacking suggest free energies measured in oligonucleotides reflect the strengths of fundamental ends) adjacent to base pairs (Sugimoto et al., 1987a; Turner et al., 1988; Burkard et al., 1999). Figure 8-17 (a) illustrates dangling end sequences in yeast phenylalanine tRNA that have ΔG values more favorable than -1 kcal mol in Table 8-5. These are strongly stacking sequences in oligonucleotides, and in the tRNA structure each such dangling end is stacked on its adjacent base pair. Figure 8-17 (b) illustrates dangling end sequences in tRNA that have ΔG° values less favorable than -0.4 kcal mol $^{-1}$. These weakly stacking sequences are not stacked on the adjacent base pairs in tRNA. In four out of five such cases, these weakly stacking sequences occur at places where there The third correlation is observed for stacking of unpaired nucleotides (dangling interactions that will help determine 3D as well as secondary structure.

by site directed mutagenesis (Michel et al., 1990; Costa and Michel, 1995), and by tions with 2'-OH groups, can be used for aligning helixes (Sugimoto et al., 1989; Pyle alignment was suggested by a phylogenetic analysis of about 80 sequences (Michel and Westhof, 1990). This analysis also suggested several other tertiary interactions such as triple helix formation and interactions with tetraloops. Some have been confirmed X-ray diffraction (Pley et al., 1994; Cate et al., 1996). Presumably, as more structures are determined, more patterns will be recognized. For example, pairing between Studies of oligonucleotides binding to natural RNAs are suggesting additional interactions that will be important for determining 3D structure. For example, interacand Cech, 1991; Bevilacqua and Tumer, 1991; Pyle et al., 1992). This type of helix complementary loops could also be important.

10. HELIX-HELIX TRANSITIONS

extensive characterization of these transitions, there is no consensus on the quantitative solvation, and steric effects are important. There is also no clearly documented case in which these types of transitions serve a known physiological function. Discovery of effective salt concentration and water activity. For example, A \rightarrow Z, B \rightarrow Z, and Thus, for example, high temperature tends to promote the Z conformation. Despite contributions of various fundamental interactions. It is likely, however, that charge, polyribonucleotides. Table 8-16 lists conditions under which various conformations are stable. Often unusual conformations are induced by conditions that affect the $B\,\rightarrow\,A$ transitions are all promoted by ethanol. High salt promotes A and B \rightarrow Z transitions. Another similarity is that these transitions typically have small ΔH^o values that are often positive (Klump and Jovin, 1987; Chaires and Sturtevant, 1986). which a helix changes conformation. Transitions between A, B, and Z forms have been studied. The Z form is most stable for alternating CG sequences, but variations are possible (Jovin and Soumpasis, 1987). The A and B forms can be stabilized for random sequence deoxy polymers, but the B form has not been observed for This chapter has focused on helix-coil transitions. Transitions are also known functional significance is probably only a matter of time.

APPENDIX

A.1 Statistical thermodynamics of transitions

The two-state model for deriving thermodynamic parameters from spectroscopic data is not general. While short oligomers often exhibit two-state transitions, long oligomers

Conditions Favoring Unusual Conformations **Table 8.16**

	Reference	Cation or Salt	Solvent) -
	(9801)	4.8-6 M NaCIO.	н,о	Š
7 + V	TUZ CI 41: (1780)	OLOWN B .	204 FrOH	ฆ
[Poly(rCrG)] (Cruz et al (1986)	TO WELL		37 00
•	Cruz et al., (1986)	S-7 M NaBr	H,O	200
	Tralson et al., (1987)	3.8 M MgCI,	н,о	გ~
	Pohl and Iovin (1972)	4 M NaCi	н,о	23
	Dobl and Iouin (1972)	2.5 M NaCIO,	H,O	ង
[rolylace)	Part (1976): Hall and Marking (1984) ~ 5 × 10 ⁻⁴ M Na	~ 5 × 10-4 M Na*	10-50% EIOH	20-50
	Fold (1970), Mail and Machine (1983)	M MoCi	н,о	22
	Fold and Selected (1972)	2 x 10-5 M Co(NH.)2+	H,O	20
	Behe and Enternfeld (1981)	> 0.7 M NaCl		~30
7+9	Delic and Colombal (1981)	> 6 × 10-4 M MgCl.	н,о	~30
B + A IDNA	[roy(dm)CdO] Belle and Classification (1974)	~ 5 × 10 * M NaCl	80% EtOH	5-30

In any statistical thermodynamic model, all the thermodynamic information is contained in the molecular partition function, q:

$$q = \sum_{j=0}^{n} \exp[-G_j/RT]$$
 (8-A.1)

possible configurations. When only duplex formation is being considered, G_j for the completely single stranded state is set at 0, and the remaining G_j values are replaced by ΔG_i values, the difference in free energy between a given duplex configuration and where G_j is the free energy of the jth configuration, and the summation is over all the single-stranded state, to give

$$q = 1 + \sum_{i=1}^{n} \exp[-\Delta G_i/RT]$$
 (8-A.2)

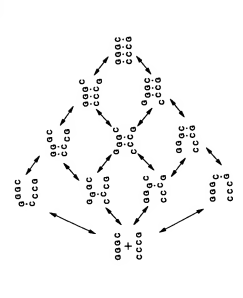
It is more convenient, however, to work with the conformational partition function, q_{ϵ} :

$$q_c = q - 1 = \sum_{i=1}^{n} \exp[-\Delta G_i/RT] = \sum_{i=1}^{n} K_i$$
 (8-A.3)

model complicated enough to describe the experimental results, but simple enough to permit unambiguous extraction of the necessary parameters from data. This approach strands. Any model can now be used to derive an expression for q_c . The trick is to use a will be illustrated with the simplest model, the zipper model (Bloomfield et al., 1974; where K_i is the equilibrium constant for forming duplex configuration i from single Cantor and Schimmel, 1980; Poland, 1978).

for a sequence such as G₃C + GC₃ (see Figure 8-18). With these assumptions, the equilibrium constant for forming each duplex configuration with j base pairs from the single strands is κs^{-1} . If there are $g_i(N)$ distinguishable duplexes with j base pairs that can be formed by a single strand with N bases, then Eq. 8-8-3. can be written In the zipper model, only one helical region is allowed per duplex. To illustrate the process in its simplest form, we assume that the equilibrium constant for adding for initiating the helix is κ (often labeled σs in the literature); and that only perfectly aligned helixes are stable enough to contribute to q. These approximations are adequate one base pair to an existing helix is always the same, s; that the equilibrium constant

$$q_c = \kappa \sum_{j=1}^{N} g_j(N) s^{j-1}$$
 (8-A.4)



Application of the zipper model to calculate the conformational partition function, $q_{\rm s}$ for duplex formation by $\rm G_{\rm s}C+GC_{\rm s}$. The zipper model allows only one helical region per duplex. Figure 8-18

 $q_c = 4\kappa + 3\kappa s + 2\kappa s^2 + \kappa s^3 = \kappa \sum_{j=1}^4 (4 \cdot j + 1) s^{j+1}$

When the two strands in the duplex have different sequence (i.e., are nonself-complementary), and only completely aligned duplexes are allowed (see Fig. 8-18), then $g_j(N) = N - j + 1$, giving

$$q_c = \kappa \sum_{j=1}^{N} (N-j+1) s^{j-1} = \kappa \left[(N+1) \sum_{j=1}^{N} s^{j-1} - \sum_{j=1}^{N} j s^{j-1} \right]$$
 (8-A.5)

These are related to the finite geometric series, for which it can be shown that

$$\sum_{i=1}^{N} s^{i-1} = \frac{s^{N} - 1}{s - 1}$$
 (8-A.6)

$$\sum_{i=1}^{N} j_i j^{-1} = \frac{N z^{N+1} - (N+1) z^N + 1}{(s-1)^2}$$
 (8-A.7)

S

$$q_r = \kappa \left(\frac{(N+1)(s^N - 1)}{s - 1} - \frac{Ns^{N+1} - (N+1)s^N + 1}{(s - 1)^2} \right) = \kappa \frac{s^{N+1} - (N+1)s + N}{(s - 1)^2}$$

Note that if s>1 and N is large, $q_c \approx \kappa s^{N-1}$. Thus, in this limit q_c is simply the equilibrium constant for the two-state model.

To determine q_{ϵ} it is necessary to measure κ and s. One way to do this is by analyzing optical melting curves. In the simplest case, the concentrations of the two complementary strands are equal, and the absorbance A of a sample depends only on the number of base pairs in the sample. For this case, the fraction X_b of bases paired is

$$X_{b} = \frac{A - A_{t}}{A_{d} - A_{t}}$$
 (8-A.9)

Here A_i and A_a are absorbances when the sample is completely single strands or fully paired duplexes, respectively. (Note the similarity to Eq. 8-14, with α corresponding to X_b .) We need to express X_b in terms of q_c . For nonself-complementary strands, A_b and B, the equilibrium can be written

$$A + B \rightleftharpoons C_1 + C_2 + \dots + C_n$$
 (8-A.10)

where C₁, C₂, ..., C_s are the possible configurations of duplex (see Fig. 8-18). Assuming the total concentrations of strands A and B are equal..

$$q_r = \frac{[C_t] + [C_2] + \dots + [C_s]}{[A_l]B_l} = \frac{0.5XC_T}{[0.5(1 - X)C_T]^2} = \frac{2X}{(1 - X)^2C_T}$$
 (8-A.11)

where C_T is the total concentration of strands,

$$C_T = \{A\} + \{B\} + 2\sum_{i=1}^{n} \{C_i\}$$
 (8-A.12)

and X is the fraction of strands in duplex,

$$X = 2(|C_1| + |C_2| + \cdots + |C_n|)/C_T$$
 (8-A.13)

The average number of base pairs per duplex, $\langle n \rangle$, can be calculated by realizing that the fraction of duplexes with j base pairs, f_i , is equal to $\kappa g_j(N) x^{-1}$, normalized by the sum over all j, q,. Thus

$$\langle n \rangle = \sum_{j=1}^{N} i j_j = \frac{1}{q_c} \sum_{j=1}^{N} j \kappa g_j(N) s^{j-1}$$

= $\frac{1}{q_c} \frac{d}{ds} \sum_{j=1}^{N} \kappa g_j(N) s^j = \frac{1}{q_c} \frac{d(sq_c)}{ds}$ (8-A.14)

This equation allows X_b to be expressed in terms of X and q_c :

$$X_{b} = \frac{X < n >}{N} = \frac{X}{Nq_{c}} \frac{d(sq_{c})}{ds}$$
 (8-A.15)

Solving Eq. 8-A.11 for X in terms of q, leads to

$$X_b = \frac{1 + q_c C_T - \sqrt{1 + 2q_c C_T}}{Nq_c^2 C_c} \frac{d(sq_c)}{ds}$$
 (8-A.16)

By using Eq. 8-A.9, X_b can be measured as a function of C_T and/or N, and the resulting simultaneous equations solved for κ and s.

each base pair (e.g., see Table 8-4). Many duplexes contain loops of various kinds so that more than one helical region is present. In this case, additional helix initiation parameters must be added that are different from κ . In general, the absorbance will not depend only on the number of base pairs (Poland, 1978). Thus X_b in Eq. 8-A.9 must be it is not general. For intramolecular helix formation with the same assumptions, the cases, however, it is more complicated. Most sequences do not have the same s for replaced by a more complicated function. Clearly, the number of parameters increases rapidly for real systems, which explains the relatively rare application of statistical While the above example illustrates the main features of the statistical approach, theory is similar (Bloomfield et al., 1974; Cantor and Schimmel, 1980). For most real

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